

Zhang 10/532690

=> d his nofile

(FILE 'HOME' ENTERED AT 10:48:36 ON 21 NOV 2006)

FILE 'HCAPLUS' ENTERED AT 10:48:46 ON 21 NOV 2006

E US2005-532690/APPS

L1 1 SEA ABB=ON PLU=ON US2005-532690/AP

E US2002-423253P/APPS

L2 1 SEA ABB=ON PLU=ON US2002-423253P/PRN

E WO2003-US34185/APPS

L3 1 SEA ABB=ON PLU=ON (WO2003-US34185/AP OR WO2003-US34185/PRN)

L*** DEL 0 S NL1-L3

L4 1 SEA ABB=ON PLU=ON (L1 OR L2 OR L3)

D SCAN

FILE 'REGISTRY' ENTERED AT 10:57:50 ON 21 NOV 2006

E 4-4-BENZYLOXY-3-CHLOROPHENYL-4-OXOBUTANOIC ACID

E 4-4-BENZYLOXY-3-CHLOROPHENYL-4-OXOBUTANOIC ACID/CN

E 4-BENZYLOXY-3-CHLOROPHENYL-4-OXOBUTANOIC ACID/CN

FILE 'HCAPLUS' ENTERED AT 10:59:07 ON 21 NOV 2006

SEL RN L4

FILE 'REGISTRY' ENTERED AT 10:59:17 ON 21 NOV 2006

L5 16 SEA ABB=ON PLU=ON (102513-61-1/BI OR 13335-57-4/BI OR
202577-82-0/BI OR 371251-24-0/BI OR 373596-81-7/BI OR 373596-82
-8/BI OR 373596-84-0/BI OR 387844-34-0/BI OR 39208-08-7/BI OR
53090-45-2/BI OR 60525-32-8/BI OR 63539-02-6/BI OR 73083-19-9/B
I OR 74362-70-2/BI OR 74362-73-5/BI OR 77513-51-0/BI)
D SCAN

L6 4 SEA ABB=ON PLU=ON L5 AND BUTANOIC
D SCAN

FILE 'REGISTRY' ENTERED AT 11:07:18 ON 21 NOV 2006

E BENZENE BUTANOIC ACID/CN

L7 1 SEA ABB=ON PLU=ON "BENZENE BUTANOIC ACID, ((3-CHLOROPHENYL) MET
HYLENE) HYDRAZIDE"/CN
D SCAN

E BENZENE BUTANOIC ACID/CN

L8 1 SEA ABB=ON PLU=ON "BENZENE BUTANOIC ACID"/CN
D SCAN

E BENZENE BUTANOIC ACID/CN

E BENZYLOXY/CN

L9 1 SEA ABB=ON PLU=ON BENZYLOXY/CN
D SCAN

FILE 'STNGUIDE' ENTERED AT 11:11:19 ON 21 NOV 2006

FILE 'REGISTRY' ENTERED AT 11:13:48 ON 21 NOV 2006

L10 STRUCTURE uploaded

L11 1 SEA SSS SAM L10

FILE 'STNGUIDE' ENTERED AT 11:14:04 ON 21 NOV 2006

FILE 'REGISTRY' ENTERED AT 11:14:29 ON 21 NOV 2006

STRUCTURE uploaded

L12 1 SEA SSS SAM L12

L13 8 SEA SSS FUL L12

D SCAN

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L15 1 SEA ABB=ON PLU=ON L5 AND L14
D SCAN
SAVE L14 NANCY690/A TEMP
L16 8 SEA ABB=ON PLU=ON (L14 OR L15)

FILE 'HCAPLUS' ENTERED AT 11:16:08 ON 21 NOV 2006

L17 4 SEA ABB=ON PLU=ON L16 (L) (THU OR PKT OR DMA OR PAC OR
BAC) /RL
D KWIC
E DIABETES/CT
E E3+ALL
E E2+ALL

L18 2178 SEA ABB=ON PLU=ON "DIABETES INSIPIDUS"/CT
L19 12931 SEA ABB=ON PLU=ON "DIABETES INSIPIDUS"+OLD/CT
E DIABETES/CT
E E3+ALL
E E3+ALL

L20 75871 SEA ABB=ON PLU=ON "DIABETES MELLITUS"/CT
E DIABETES/CT
E E4+ALL

L21 12931 SEA ABB=ON PLU=ON "DIABETES INSIPIDUS"+OLD/CT
E DIABETES/CT
E E7+ALL

L22 297 SEA ABB=ON PLU=ON "DIABETES INSIPIDUS (L) NEPHROGENIC"/CT
L23 117864 SEA ABB=ON PLU=ON ?DIABETES?
E ATHEROSCLEROSIS/CT
E E3+ALL

L24 37882 SEA ABB=ON PLU=ON ATHEROSCLEROSIS+OLD/CT
E ATHEROSCLEROSIS/CT
E ARTERIOSCLEROSIS/CT
E E3+ALL

L25 41547 SEA ABB=ON PLU=ON ARTERIOSCLEROSIS+NT/CT
E ARTERIOSCLEROSIS/CT
E E4+ALL
E OBESITY/CT
E E3+ALL

L26 28762 SEA ABB=ON PLU=ON OBESITY+NT/CT
E HYPERTENSION/CT
E E3+ALL

L27 51377 SEA ABB=ON PLU=ON HYPERTENSION/CT
E FATTY LIVER DISEASE/CT
E E3+ALL
E E2+ALL

L28 11032 SEA ABB=ON PLU=ON "LIVER, DISEASE (L) FATTY"+OLD/CT
E NEPHROPATHY/CT
E E3+ALL
E E2+ALL

L29 40906 SEA ABB=ON PLU=ON "KIDNEY, DISEASE"+OLD+NT/CT
L30 68189 SEA ABB=ON PLU=ON "KIDNEY, DISEASE"+OLD, NT/CT
E RETINOPATHY/CT
E E3+ALL
E E2+ALL

L31 7915 SEA ABB=ON PLU=ON "EYE, DISEASE (L) RETINOPATHY"+OLD/CT
E FOOT ULCERATION/CT
E FOOT /CT
E E+ALL
E E3+ALL
E FOOT/CT
E E3+ALL

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E ULCERATION/CT
E CATARACT/CT
E E3+ALL
L32 5862 SEA ABB=ON PLU=ON CATARACT+OLD/CT
E CATARACT/CT
E E4+ALL
E HYPERLIPIDEMIA/CT
E E3+ALL
L33 11861 SEA ABB=ON PLU=ON HYPERLIPIDEMIA+OLD, NT/CT
E CACHEXIA/CT
E E3+ALL
L34 2566 SEA ABB=ON PLU=ON CACHEXIA/CT
L35 234979 SEA ABB=ON PLU=ON FATTY LIVER DISEASE? OR FOOT ULCER? OR
FEET ULCER? OR INSULIN RESISTANCE? OR OBESITY? OR HYPERLIPIDEMI
? OR ATHEROSCLERO? OR ARTERIOSCLER? OR HYPERTENS? OR NEPHROPATH
? OR NEUROPATH? OR RETINOPATH? OR CACHEXIA
L36 381179 SEA ABB=ON PLU=ON (L18 OR L19 OR L20 OR L21 OR L22 OR L23 OR
L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31 OR L32 OR
L33 OR L34 OR L35)
L37 3 SEA ABB=ON PLU=ON L36 AND L17
L38 5 SEA ABB=ON PLU=ON L16 AND L36
L39 5 SEA ABB=ON PLU=ON (L37 OR L38)
D KWIC
L40 4 SEA ABB=ON PLU=ON L39 NOT L4
E HODGE K/AU
L41 12 SEA ABB=ON PLU=ON ("HODGE K"/AU OR "HODGE KIRVIN L"/AU)
E SHARMA S/AU
L42 3442 SEA ABB=ON PLU=ON ("SHARMA S"/AU OR "SHARMA S A"/AU OR
"SHARMA S A N"/AU OR "SHARMA S AMITA"/AU OR "SHARMA S B"/AU OR
"SHARMA S C"/AU OR "SHARMA S C L"/AU OR "SHARMA S CHIDANANDA"/AU
OR "SHARMA S D"/AU OR "SHARMA S D GURUMAYUM"/AU OR "SHARMA S
DAS"/AU OR "SHARMA S G"/AU OR "SHARMA S H K"/AU OR "SHARMA S
J"/AU OR "SHARMA S K"/AU OR "SHARMA S KUMAR"/AU OR "SHARMA S
L"/AU OR "SHARMA S M"/AU OR "SHARMA S N"/AU OR "SHARMA S P"/AU
OR "SHARMA S R"/AU OR "SHARMA S RAMA GOPAL"/AU OR "SHARMA S
S"/AU OR "SHARMA S SEN"/AU OR "SHARMA S SHELLEY"/AU OR "SHARMA
S SHELLY"/AU OR "SHARMA S V"/AU OR "SHARMA SHALINI"/AU)
E VON BORSTEL R/AU
L43 114 SEA ABB=ON PLU=ON ("VON BORSTEL R"/AU OR "VON BORSTEL R
C"/AU OR "VON BORSTEL REID"/AU OR "VON BORSTEL REID W"/AU OR
"VON BORSTEL REID WARREN"/AU)
E VONBORSTEL R/AU
L44 2 SEA ABB=ON PLU=ON "VONBORSTEL REID W"/AU
L45 116 SEA ABB=ON PLU=ON (L43 OR L44)
E WOLPE S/AU
L46 33 SEA ABB=ON PLU=ON ("WOLPE S"/AU OR "WOLPE S D"/AU OR "WOLPE
STEPHEN"/AU OR "WOLPE STEPHEN D"/AU OR "WOLPE STEVE D"/AU OR
"WOLPE STEVEN"/AU)
L47 7 SEA ABB=ON PLU=ON (L41 AND (L42 OR L45 OR L46)) OR (L42 AND
(L45 OR L46)) OR (L45 AND L46)
L48 4 SEA ABB=ON PLU=ON L40 NOT L47

FILE 'MEDLINE' ENTERED AT 11:28:39 ON 21 NOV 2006

FILE 'HCAPLUS' ENTERED AT 11:29:02 ON 21 NOV 2006

L49 6 SEA ABB=ON PLU=ON (L17 OR L48)
L50 5 SEA ABB=ON PLU=ON L49 NOT (L47 OR L4)

FILE 'MEDLINE' ENTERED AT 11:29:39 ON 21 NOV 2006

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L51 0 SEA ABB=ON PLU=ON L16
L52 FILE 'EMBASE, BIOSIS, CAOLD' ENTERED AT 11:29:56 ON 21 NOV 2006
L52 0 SEA ABB=ON PLU=ON L16
L53 FILE 'WPIX' ENTERED AT 11:30:02 ON 21 NOV 2006
L53 1 SEA SSS FUL L12
L54 FILE 'REGISTRY' ENTERED AT 11:30:42 ON 21 NOV 2006
L54 D BROWSE L15
L54 0 SEA ABB=ON PLU=ON 74362-73-5/CRN
L55 FILE 'HCAPLUS' ENTERED AT 11:31:18 ON 21 NOV 2006
L55 2 SEA ABB=ON PLU=ON L15
L56 7 SEA ABB=ON PLU=ON (L55 OR L50)
L57 6 SEA ABB=ON PLU=ON L56 NOT (L47 OR L4)
L58 FILE 'MEDLINE, EMBASE, BIOSIS, CAOLD' ENTERED AT 11:31:55 ON 21 NOV 2006
L58 0 SEA ABB=ON PLU=ON L15
L59 FILE 'WPIX' ENTERED AT 11:32:13 ON 21 NOV 2006
L59 FILE 'REGISTRY' ENTERED AT 11:32:24 ON 21 NOV 2006
L59 D BROWSE L15
L59 FILE 'WPIX' ENTERED AT 11:32:39 ON 21 NOV 2006
L59 0 SEA ABB=ON PLU=ON BENZENE BUTANOIC ACID/CN
L59 E BENZENE BUTANOIC ACID, 3-CHLORO-Γ-OXO-4-(PHENYLMETHOXY)-
L*** DEL FILE 'HCAPLUS' ENTERED AT 11:33:15 ON 21 NOV 2006
L60 1 S L1
L60 8 SEA ABB=ON PLU=ON L16
L61 1 SEA ABB=ON PLU=ON L60 NOT (L57 OR L47 OR L1)
L62 1 SEA ABB=ON PLU=ON L60 NOT (L57 OR L47 OR L4)
L62 D KWIC
L63 7 SEA ABB=ON PLU=ON (L61 OR L62 OR L57)
L64 0 SEA ABB=ON PLU=ON L60 NOT (L63 OR L47 OR L4)
L65 FILE 'HCAPLUS' ENTERED AT 11:34:17 ON 21 NOV 2006
L65 FILE 'WPIX' ENTERED AT 11:34:23 ON 21 NOV 2006
L65 D QUE L47
L65 D QUE L63
L65 D QUE L53
L65 FILE 'STNGUIDE' ENTERED AT 11:34:37 ON 21 NOV 2006
L65 FILE 'HCAPLUS, WPIX' ENTERED AT 11:34:47 ON 21 NOV 2006
L65 15 DUP REM L47 L63 L53 (0 DUPLICATES REMOVED)
L65 ANSWERS '1-14' FROM FILE HCAPLUS
L65 ANSWER '15' FROM FILE WPIX
L66 FILE 'WPIX' ENTERED AT 11:35:04 ON 21 NOV 2006
L66 1 SEA ABB=ON PLU=ON L53/DCR
L66 SEL SDCN L53
L67 EDIT E1 SDCN DCN
L67 1 SEA ABB=ON PLU=ON RAECKI/DCN
L67 SEL DCSE L53
L67 EDIT E2 DCSE DCRE

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L68 0 SEA ABB=ON PLU=ON 905973-0-0-0/DCRE
L69 1 SEA ABB=ON PLU=ON (L67 OR L68)

FILE 'HCAPLUS' ENTERED AT 11:36:48 ON 21 NOV 2006

FILE 'WPIX' ENTERED AT 11:36:53 ON 21 NOV 2006

 D QUE L47
 D QUE L63
 D QUE L53
 D QUE L69

FILE 'STNGUIDE' ENTERED AT 11:37:03 ON 21 NOV 2006

FILE 'HCAPLUS, WPIX' ENTERED AT 11:37:12 ON 21 NOV 2006
L70 15 DUP REM L47 L63 L53 L69 (1 DUPLICATE REMOVED)
 ANSWERS '1-14' FROM FILE HCAPLUS
 ANSWER '15' FROM FILE WPIX
 D IBIB ABS HITIND HITSTR RETABLE L70 1-14
 D IDE L53 TOT
 D ALL ABEQ TECH L70 TOT

FILE 'WPIX' ENTERED AT 11:38:52 ON 21 NOV 2006

 D ALL ABEQ TECH L69 TOT

FILE 'REGISTRY' ENTERED AT 12:02:33 ON 21 NOV 2006
 D BROWSE L15

FILE 'REGISTRY' ENTERED AT 12:02:51 ON 21 NOV 2006
L71 STR 74362-73-5
L72 1 SEA FAM FUL L71
 D SCAN
L73 STRUCTURE UPLOADED
L74 1 SEA SSS SAM L73
L75 8 SEA SSS FUL L73
L76 8 SEA ABB=ON PLU=ON (L75 OR L14)

=> file hcaplus

FILE 'HCAPLUS' ENTERED AT 11:36:48 ON 21 NOV 2006
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE COVERS 1907 - 21 Nov 2006 VOL 145 ISS 22
FILE LAST UPDATED: 20 Nov 2006 (20061120/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> file wpix
FILE 'WPIX' ENTERED AT 11:36:53 ON 21 NOV 2006
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FILE LAST UPDATED: 20 NOV 2006 <20061120/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200674 <200674/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

>>> YOU ARE IN THE NEW AND ENHANCED DERWENT WORLD PATENTS INDEX <<<

>>> FOR DETAILS ON THE NEW AND ENHANCED DERWENT WORLD PATENTS INDEX PLEASE VISIT:
[<<<](http://www.stn-international.de/stndatabases/details/dwpi_r.html)

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<http://scientific.thomson.com/support/patents/coverage/latestupdates/>

PLEASE BE AWARE OF THE NEW IPC REFORM IN 2006, SEE
http://www.stn-international.de/stndatabases/details/ ipc_reform.html and
<http://scientific.thomson.com/media/scpdf/ ipcrdwpi.pdf>

>>> FOR DETAILS ON THE NEW AND ENHANCED DERWENT WORLD PATENTS INDEX PLEASE SEE
[<<<](http://www.stn-international.de/stndatabases/details/dwpi_r.html)

>>> YOU ARE IN THE NEW AND ENHANCED DERWENT WORLD PATENTS INDEX <<<
'BIX' IS DEFAULT SEARCH FIELD FOR 'WPIX' FILE

=> d que 147

L41 12 SEA FILE=HCAPLUS ABB=ON PLU=ON ("HODGE K"/AU OR "HODGE KIRVIN L"/AU)

L42 3442 SEA FILE=HCAPLUS ABB=ON PLU=ON ("SHARMA S"/AU OR "SHARMA S A"/AU OR "SHARMA S A N"/AU OR "SHARMA S AMITA"/AU OR "SHARMA S B"/AU OR "SHARMA S C"/AU OR "SHARMA S C L"/AU OR "SHARMA S CHIDANANDA"/AU OR "SHARMA S D"/AU OR "SHARMA S D GURUMAYUM"/AU OR "SHARMA S DAS"/AU OR "SHARMA S G"/AU OR "SHARMA S H K"/AU OR "SHARMA S J"/AU OR "SHARMA S K"/AU OR "SHARMA S KUMAR"/AU OR "SHARMA S L"/AU OR "SHARMA S M"/AU OR "SHARMA S N"/AU OR "SHARMA S P"/AU OR "SHARMA S R"/AU OR "SHARMA S RAMA GOPAL"/AU OR "SHARMA S S"/AU OR "SHARMA S SEN"/AU OR "SHARMA S SHELLEY"/AU OR "SHARMA S SHELLY"/AU OR "SHARMA S V"/AU OR "SHARMA SHALINI"/AU)

L43 114 SEA FILE=HCAPLUS ABB=ON PLU=ON ("VON BORSTEL R"/AU OR "VON BORSTEL R C"/AU OR "VON BORSTEL REID"/AU OR "VON BORSTEL REID W"/AU OR "VON BORSTEL REID WARREN"/AU)

L44 2 SEA FILE=HCAPLUS ABB=ON PLU=ON "VONBORSTEL REID W"/AU

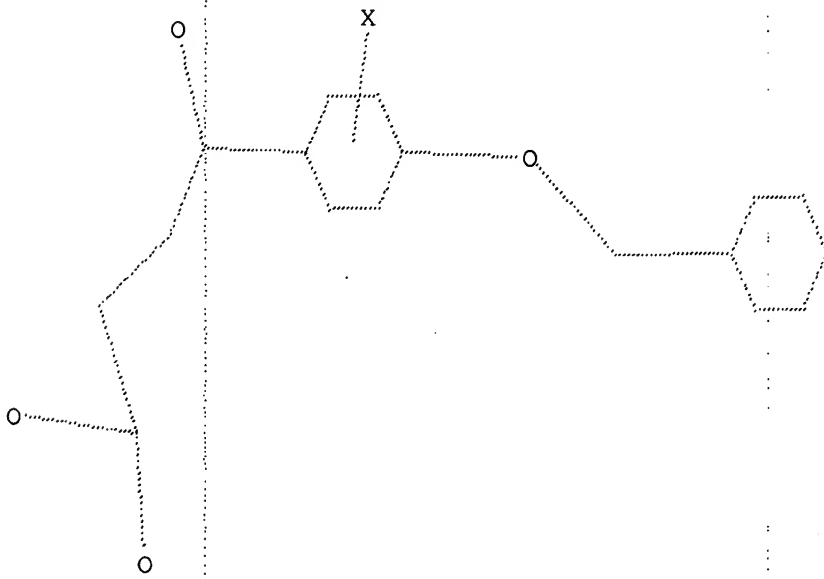
L45 116 SEA FILE=HCAPLUS ABB=ON PLU=ON (L43 OR L44)

L46 33 SEA FILE=HCAPLUS ABB=ON PLU=ON ("WOLPE S"/AU OR "WOLPE S D"/AU OR "WOLPE STEPHEN"/AU OR "WOLPE STEPHEN D"/AU OR "WOLPE

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STEVE D"/AU OR "WOLPE STEVEN"/AU)
L47 7 SEA FILE=HCAPLUS ABB=ON PLU=ON (L41 AND (L42 OR L45 OR L46))
OR (L42 AND (L45 OR L46)) OR (L45 AND L46)

=> d que 163
L1 1 SEA FILE=HCAPLUS ABB=ON PLU=ON US2005-532690/AP
L2 1 SEA FILE=HCAPLUS ABB=ON PLU=ON US2002-423253P/PRN
L3 1 SEA FILE=HCAPLUS ABB=ON PLU=ON (WO2003-US34185/AP OR
WO2003-US34185/PRN)
L4 1 SEA FILE=HCAPLUS ABB=ON PLU=ON (L1 OR L2 OR L3)
L5 16 SEA FILE=REGISTRY ABB=ON PLU=ON (102513-61-1/BI OR 13335-57-4
/BI OR 202577-82-0/BI OR 371251-24-0/BI OR 373596-81-7/BI OR
373596-82-8/BI OR 373596-84-0/BI OR 387844-34-0/BI OR 39208-08-
7/BI OR 53090-45-2/BI OR 60525-32-8/BI OR 63539-02-6/BI OR
73083-19-9/BI OR 74362-70-2/BI OR 74362-73-5/BI OR 77513-51-0/B
I)
L12 STR



Structure attributes must be viewed using STN Express query preparation.

L14 8 SEA FILE=REGISTRY SSS FUL L12
L15 1 SEA FILE=REGISTRY ABB=ON PLU=ON L5 AND L14
L16 8 SEA FILE=REGISTRY ABB=ON PLU=ON (L14 OR L15)
L17 4 SEA FILE=HCAPLUS ABB=ON PLU=ON L16 (L) (THU OR PKT OR DMA OR
PAC OR BAC)/RL
L18 2178 SEA FILE=HCAPLUS ABB=ON PLU=ON "DIABETES INSIPIDUS"/CT
L19 12931 SEA FILE=HCAPLUS ABB=ON PLU=ON "DIABETES INSIPIDUS"+OLD/CT
L20 75871 SEA FILE=HCAPLUS ABB=ON PLU=ON "DIABETES MELLITUS"/CT
L21 12931 SEA FILE=HCAPLUS ABB=ON PLU=ON "DIABETES INSIPIDUS"+OLD/CT
L22 297 SEA FILE=HCAPLUS ABB=ON PLU=ON "DIABETES INSIPIDUS (L)
NEPHROGENIC"/CT
L23 117864 SEA FILE=HCAPLUS ABB=ON PLU=ON ?DIABETES?
L24 37882 SEA FILE=HCAPLUS ABB=ON PLU=ON ATHEROSCLEROSIS+OLD/CT
L25 41547 SEA FILE=HCAPLUS ABB=ON PLU=ON ARTERIOSCLEROSIS+NT/CT

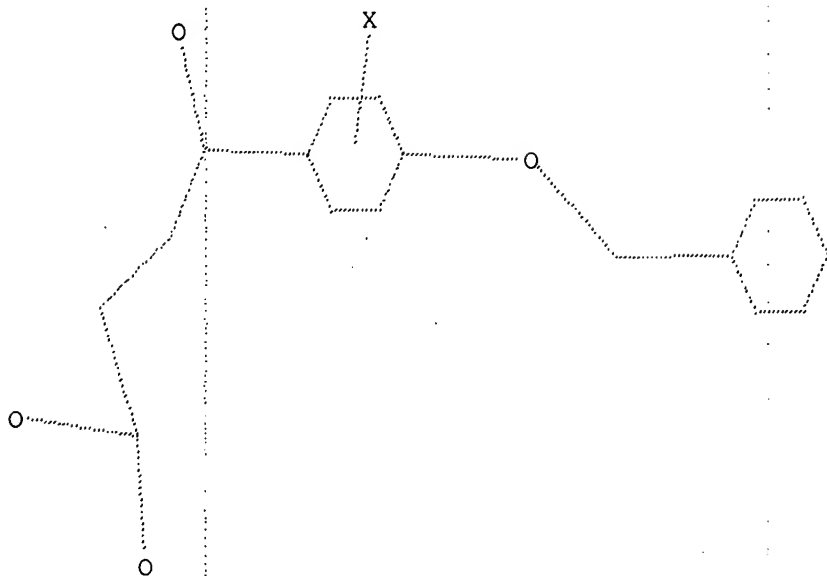
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L26 28762 SEA FILE=HCAPLUS ABB=ON PLU=ON OBESITY+NT/CT
L27 51377 SEA FILE=HCAPLUS ABB=ON PLU=ON HYPERTENSION/CT
L28 11032 SEA FILE=HCAPLUS ABB=ON PLU=ON "LIVER, DISEASE (L) FATTY"+OLD
/CT
L29 40906 SEA FILE=HCAPLUS ABB=ON PLU=ON "KIDNEY, DISEASE"+OLD+NT/CT
L30 68189 SEA FILE=HCAPLUS ABB=ON PLU=ON "KIDNEY, DISEASE"+OLD,NT/CT
L31 7915 SEA FILE=HCAPLUS ABB=ON PLU=ON "EYE, DISEASE (L) RETINOPATHY"
+OLD/CT
L32 5862 SEA FILE=HCAPLUS ABB=ON PLU=ON CATARACT+OLD/CT
L33 11861 SEA FILE=HCAPLUS ABB=ON PLU=ON HYPERLIPIDEMIA+OLD,NT/CT
L34 2566 SEA FILE=HCAPLUS ABB=ON PLU=ON CACHEXIA/CT
L35 234979 SEA FILE=HCAPLUS ABB=ON PLU=ON FATTY LIVER DISEASE? OR FOOT
ULCER? OR FEET ULCER? OR INSULIN RESISTANCE? OR OBESITY? OR
HYPERLIPIDEMI? OR ATHEROSCLERO? OR ARTERIOSCLER? OR HYPERTENS?
OR NEPHROPATH? OR NEUROPATH? OR RETINOPATH? OR CACHEXIA
L36 381179 SEA FILE=HCAPLUS ABB=ON PLU=ON (L18 OR L19 OR L20 OR L21 OR
L22 OR L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR
L31 OR L32 OR L33 OR L34 OR L35)
L37 3 SEA FILE=HCAPLUS ABB=ON PLU=ON L36 AND L17
L38 5 SEA FILE=HCAPLUS ABB=ON PLU=ON L16 AND L36
L39 5 SEA FILE=HCAPLUS ABB=ON PLU=ON (L37 OR L38)
L40 4 SEA FILE=HCAPLUS ABB=ON PLU=ON L39 NOT L4
L41 12 SEA FILE=HCAPLUS ABB=ON PLU=ON ("HODGE K"/AU OR "HODGE
KIRVIN L"/AU)
L42 3442 SEA FILE=HCAPLUS ABB=ON PLU=ON ("SHARMA S"/AU OR "SHARMA S
A"/AU OR "SHARMA S A N"/AU OR "SHARMA S AMITA"/AU OR "SHARMA S
B"/AU OR "SHARMA S C"/AU OR "SHARMA S C L"/AU OR "SHARMA S
CHIDANANDA"/AU OR "SHARMA S D"/AU OR "SHARMA S D GURUMAYUM"/AU
OR "SHARMA S DAS"/AU OR "SHARMA S G"/AU OR "SHARMA S H K"/AU
OR "SHARMA S J"/AU OR "SHARMA S K"/AU OR "SHARMA S KUMAR"/AU
OR "SHARMA S L"/AU OR "SHARMA S M"/AU OR "SHARMA S N"/AU OR
"SHARMA S P"/AU OR "SHARMA S R"/AU OR "SHARMA S RAMA GOPAL"/AU
OR "SHARMA S S"/AU OR "SHARMA S SEN"/AU OR "SHARMA S SHELLY"/AU
OR "SHARMA S SHELLY"/AU OR "SHARMA S V"/AU OR "SHARMA
SHALINI"/AU)
L43 114 SEA FILE=HCAPLUS ABB=ON PLU=ON ("VON BORSTEL R"/AU OR "VON
BORSTEL R C"/AU OR "VON BORSTEL REID"/AU OR "VON BORSTEL REID
W"/AU OR "VON BORSTEL REID WARREN"/AU)
L44 2 SEA FILE=HCAPLUS ABB=ON PLU=ON "VONBORSTEL REID W"/AU
L45 116 SEA FILE=HCAPLUS ABB=ON PLU=ON (L43 OR L44)
L46 33 SEA FILE=HCAPLUS ABB=ON PLU=ON ("WOLPE S"/AU OR "WOLPE S
D"/AU OR "WOLPE STEPHEN"/AU OR "WOLPE STEPHEN D"/AU OR "WOLPE
STEVE D"/AU OR "WOLPE STEVEN"/AU)
L47 7 SEA FILE=HCAPLUS ABB=ON PLU=ON (L41 AND (L42 OR L45 OR L46))
OR (L42 AND (L45 OR L46)) OR (L45 AND L46)
L48 4 SEA FILE=HCAPLUS ABB=ON PLU=ON L40 NOT L47
L49 6 SEA FILE=HCAPLUS ABB=ON PLU=ON (L17 OR L48)
L50 5 SEA FILE=HCAPLUS ABB=ON PLU=ON L49 NOT (L47 OR L4)
L55 2 SEA FILE=HCAPLUS ABB=ON PLU=ON L15
L56 7 SEA FILE=HCAPLUS ABB=ON PLU=ON (L55 OR L50)
L57 6 SEA FILE=HCAPLUS ABB=ON PLU=ON L56 NOT (L47 OR L4)
L60 8 SEA FILE=HCAPLUS ABB=ON PLU=ON L16
L61 1 SEA FILE=HCAPLUS ABB=ON PLU=ON L60 NOT (L57 OR L47 OR L1)
L62 1 SEA FILE=HCAPLUS ABB=ON PLU=ON L60 NOT (L57 OR L47 OR L4)
L63 7 SEA FILE=HCAPLUS ABB=ON PLU=ON (L61 OR L62 OR L57)

=> d que 153

L12 STR

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Structure attributes must be viewed using STN Express query preparation.
L53 1 SEA FILE=WPIX SSS FUL L12

=> d que 169
L67 1 SEA FILE=WPIX ABB=ON PLU=ON RAECKI/DCN
L68 0 SEA FILE=WPIX ABB=ON PLU=ON 905973-0-0-0/DCRE
L69 1 SEA FILE=WPIX ABB=ON PLU=ON (L67 OR L68)

=> file stnguide
FILE 'STNGUIDE' ENTERED AT 11:37:03 ON 21 NOV 2006
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AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Nov 17, 2006 (20061117/UP).

=> dup rem 147,163,153,169
FILE 'HCAPLUS' ENTERED AT 11:37:12 ON 21 NOV 2006
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PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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FILE 'WPIX' ENTERED AT 11:37:12 ON 21 NOV 2006
COPYRIGHT (C) 2006 THE THOMSON CORPORATION
PROCESSING COMPLETED FOR L47
PROCESSING COMPLETED FOR L63
PROCESSING COMPLETED FOR L53
PROCESSING COMPLETED FOR L69
L70 15 DUP REM L47 L63 L53 L69 (1 DUPLICATE REMOVED)
 ANSWERS '1-14' FROM FILE HCAPLUS

Zhang 10/532690

ANSWER '15' FROM FILE WPIX

=> d ibib abs hitind hitstr retable 170 1-14;d ide 153 tot ;d all abeq tech 170 tot

L70 ANSWER 1 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1
ACCESSION NUMBER: 2004:412751 HCAPLUS <<LOGINID::20061121>>
DOCUMENT NUMBER: 140:400084
TITLE: Oxocarboxylic acids and esters thereof for the treatment of metabolic disorders
INVENTOR(S): *Hodge, Kirvin L.; Sharma, Shalini; Von Borstel, Reid W.; Wolpe, Stephen D.*
PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA; Von Borstel, Reid W.
SOURCE: PCT Int. Appl., 22 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004041165	A2	20040521	WO 2003-US34185	20031028
WO 2004041165	A3	20050203		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2502297	AA	20040521	CA 2003-2502297	20031028
AU 2003286728	A1	20040607	AU 2003-286728	20031028
EP 1556085	A2	20050727	EP 2003-777939	20031028
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				
JP 2006507303	T2	20060302	JP 2004-550151	20031028
US 2006035970	A1	20060216	US 2005-532690	20050426
PRIORITY APPLN. INFO.:			US 2002-423253P	P 20021101
			WO 2003-US34185	W 20031028

AB Oxocarboxylic acids and esters thereof are disclosed which are useful for the treatment of various metabolic disorders, e.g. insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis.

IC ICM A61K

CC 1-10 (Pharmacology)

L70 ANSWER 2 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:177884 HCAPLUS <<LOGINID::20061121>>
DOCUMENT NUMBER: 142:279944
TITLE: Preparation of phenyl thioethers for the treatment of metabolic disorders
INVENTOR(S): *Sharma, Shalini; Von Borstel, Reid W.; Hodge, Kirvin L.*
PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA

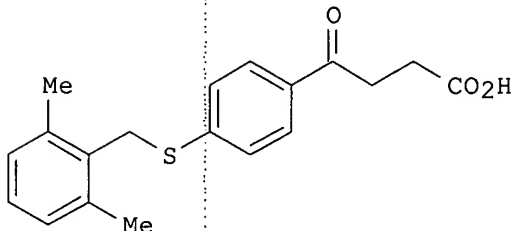
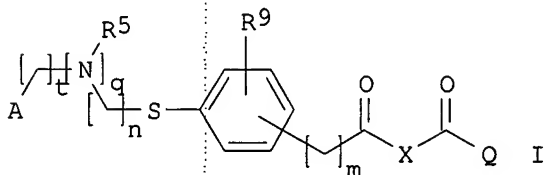
Zhang 10/532690

SOURCE: PCT Int. Appl., 42 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005018628	A1	20050303	WO 2004-US26561	20040816
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2004266673	A1	20050303	AU 2004-266673	20040816
CA 2533890	AA	20050303	CA 2004-2533890	20040816
EP 1656127	A1	20060517	EP 2004-781277	20040816
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK				
CN 1835743	A	20060920	CN 2004-80023552	20040816
NO 2006000502	A	20060503	NO 2006-502	20060131
PRIORITY APPLN. INFO.:			US 2003-496533P	P 20030820
			WO 2004-US26561	W 20040816

OTHER SOURCE(S): CASREACT 142:279944; MARPAT 142:279944

GI



AB The title compds. I [n = 1-2; m, q, t = 0-1; R5 = alkyl; R9 = H, halo,

alkyl, alkoxy; A = (un)substituted Ph, cycloalkyl, 5-6 membered heteroarom. ring having 1 or 2 ring heteroatoms selected from N, S and O and the heteroarom. ring is covalently bound to the remainder of the compound I by a ring carbon; X = CH₂; Q = OR₁ and R₁ = Me, Et; or X = CH₂CR₁₂R₁₃ or CH₂CH(NHAc) (wherein R₁₂, R₁₃ = H, Me), Q = OR₁ and R₁ = H, alkyl; or X = CH₂CH₂ and Q = NR₁₀R₁₁ (wherein one of R₁₀ and R₁₁ = H, alkyl or OH, and the other = H); alternatively, when R₁ = H, the biol. active agent can be a pharmaceutically acceptable salt of the compound I], useful for the treatment of various metabolic disorders, such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis are disclosed. E.g., a multi-step synthesis of II, starting from 2,6-dimethylbenzyl alc., was given. The pharmaceutical composition comprising the compound I is also disclosed.

IC ICM A61K031-19

ICS A61K031-235; C07C323-00

CC 25-10 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)

Section cross-reference(s): 1, 63

RETABLE

Referenced Author (RAU)	Year (R PY)	VOL (R VL)	PG (R PG)	Referenced Work (RWK)	Referenced File
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Sharma | 2002 | | WO 02100341 A2 | HCAPLUS

L70 ANSWER 3 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:962218 HCAPLUS <>LOGINID: 20061121>>

DOCUMENT NUMBER: 143:266913

TITLE: Preparation of 3-pyrazolecarboxamide derivatives as CB1 receptor modulators for the treatment of **obesity** and other diseasesINVENTOR(S): Cheng, Leifeng; Lindstedt-Alstermark, Eva-Lotte;
Boije, Anna Maria Persdotter

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 67 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

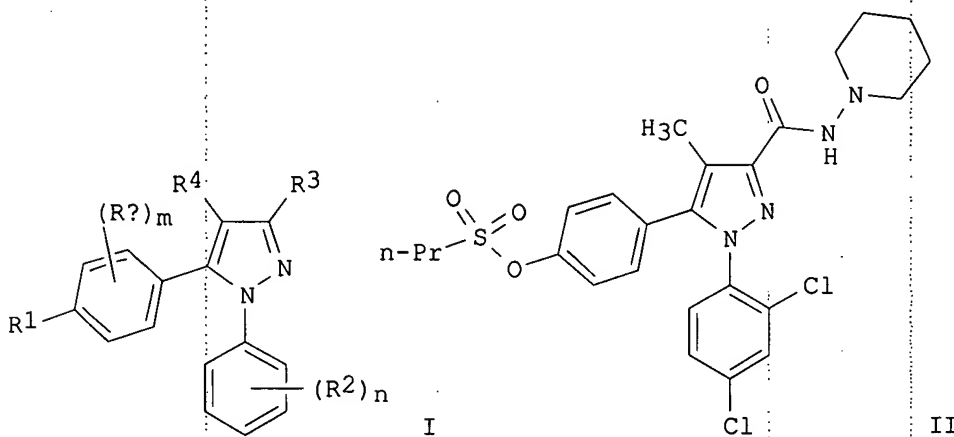
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005080343	A2	20050901	WO 2005-GB534	20050216
WO 2005080343	A3	20060112		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, SM			
RW:	BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
AU 2005214130	A1	20050901	AU 2005-214130	20050216
CA 2555331	AA	20050901	CA 2005-2555331	20050216
EP 1718617	A2	20061108	EP 2005-717730	20050216
R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,			

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, BA, HR, IS, YU				
NO 2006003787	A	20060919	NO 2006-3787	20060824
PRIORITY APPLN. INFO.:			GB 2004-3779	A 20040220
			GB 2004-20780	A 20040918
			WO 2005-GB534	W 20050216

OTHER SOURCE(S) : MARPAT 143:266913
GI



AB The title compds. I [R1 = substituted alkoxy, sulfonyl, sulfonamide or silanyl; Ra = halo, alkyl or alkoxy; m, n = 0-3; R2 = alkyl, alkoxy, etc.; R3 = substituted aminocarbonyl, etc.; R4 = H, alkyl, etc., with two exclusions, and pharmaceutically acceptable salts thereof] were prepared as CB1 receptor modulators. As an example, II was synthesized via sulfonylation of the corresponding phenol (preparation given) with n-PrSO₂Cl in 49% yield. I are active in the CB1 receptor with IC₅₀ values of < 1 μM (IC₅₀ = 6 nM for II) and believed to be selective CB1 antagonists or inverse agonists. Therefore, I and their pharmaceutical compns. may be used in the treatment of **obesity**, psychiatric disorders, neurol. disorders and so on.

IC ICM C07D231-00

CC 28-8 (Heterocyclic Compounds (More Than One Hetero Atom))

Section cross-reference(s): 1, 63

ST pyrazolecarboxamide prep ***obesity*** psychiatric neural disorder treatment; CB1 receptor modulator pyrazole carboxamide prep

IT Nervous system, disease

(Huntington's chorea, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)

IT Mental and behavioral disorders

(attention deficit disorder, treatment of; preparation of pyrazolecarboxamides CB₁ receptor modulators for treatment of **obesity** and other diseases)

IT Mental and behavioral disorders

(bipolar disorder, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other

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diseases)
IT Mental and behavioral disorders
(dementia, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Mental and behavioral disorders
(depression, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Mental and behavioral disorders
(obsession-compulsion, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Anti-Alzheimer's agents
Anticonvulsants
Antidepressants
Antibesity agents
Antiparkinsonian agents
Antipsychotics
Anxiolytics
Cardiovascular agents
Cognition enhancers
Immunomodulators
Nervous system agents
(preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Mental and behavioral disorders
(psychosis, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Shock (circulatory collapse)
(septic, treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Alzheimer's disease
Anorexia
Anxiety
Cardiovascular system, disease
Cognitive disorders
Digestive tract, disease
Drug dependence
Endocrine system, disease
Epilepsy
Immune disease
Memory disorders
Mental and behavioral disorders
Nervous system, disease
Obesity
Parkinson's disease
Reproduction disorders
Respiratory system, disease
Schizophrenia
(treatment of; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT Cannabinoid receptors
RL: BSU (Biological study, unclassified); BIOL (Biological study)
(type CB1, modulator; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)
IT 863639-64-9P 863639-65-0P
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
(drug candidate; preparation of pyrazolecarboxamides CB1 receptor modulators

for treatment of **obesity** and other diseases)

IT 863639-38-7P 863639-39-8P 863639-40-1P 863639-41-2P 863639-43-4P
 863639-44-5P 863639-46-7P 863639-52-5P 863639-56-9P 863639-58-1P
 863639-61-6P 863639-62-7P 863639-63-8P 863639-66-1P 863639-67-2P
 863639-68-3P 863639-69-4P 863639-72-9P 863639-73-0P 863639-78-5P
 863639-79-6P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)

IT 52-52-8, 1-Aminocyclopentanecarboxylic acid 70-70-2 95-92-1, Oxalic acid diethyl ester 100-39-0, Benzyl bromide 109-61-5, Propyl chloroformate 110-73-6, 2-(Ethylamino)ethanol 141-97-9, Ethyl acetoacetate 443-93-6 461-17-6, 1-Iodo-4,4,4-trifluorobutane 554-00-7, 2,4-Dichloroaniline 2213-43-6, 1-Aminopiperidine 2386-60-9, Butanesulfonyl chloride 2766-74-7, 5-Chlorothiophène-2-sulfonyl chloride 4319-49-7, 4-Aminomorpholine 10147-36-1, 1-Propanesulfonyl chloride 10307-18-3 13123-92-7, (2,4-Dichlorophenyl)hydrazine 16133-25-8, 3-Pyridinesulfonyl chloride 16629-19-9, 2-Thiophenesulfonyl chloride 18742-02-4, 2-(2-Bromoethyl)-1,3-dioxolane 22795-37-5, 3-Methylbutane-1-sulfonyl chloride 54696-05-8, 1-(4-Benzylxyphenyl)ethanone 63234-70-8, 1-Aminopiperidine hydrochloride 74784-70-6, 5-(Trifluoromethyl)pyridin-2-amine 178374-78-2 212190-25-5 845866-80-0 863639-75-2

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)

IT 4495-66-3P 35081-45-9P 57696-12-5P 60421-23-0P, Methyl 1-aminocyclopentanecarboxylate hydrochloride 152192-95-5P 178374-92-0P 178374-93-1P 502486-92-2P 503270-34-6P 863639-35-4P 863639-36-5P 863639-37-6P 863639-42-3P **863639-47-8P** 863639-48-9P 863639-49-0P 863639-50-3P 863639-51-4P 863639-53-6P 863639-54-7P 863639-55-8P 863639-57-0P 863639-59-2P 863639-60-5P 863639-70-7P 863639-71-8P 863639-74-1P 863639-76-3P 863639-77-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)

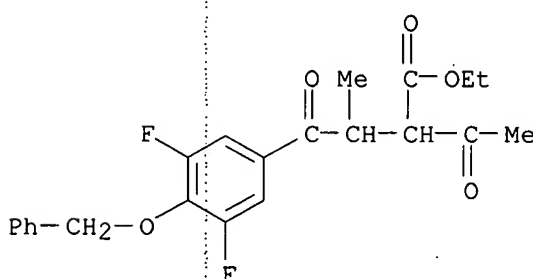
863639-47-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of pyrazolecarboxamides CB1 receptor modulators for treatment of **obesity** and other diseases)

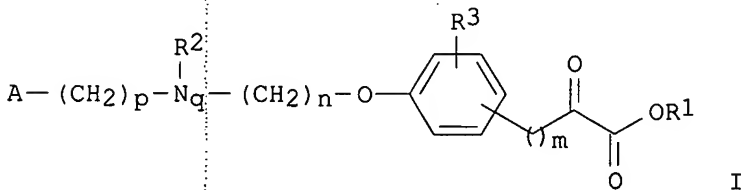
RN 863639-47-8 HCPLUS

CN Benzenebutanoic acid, α -acetyl-3,5-difluoro- β -methyl- γ -oxo-4-(phenylmethoxy)-, ethyl ester (9CI) (CA INDEX NAME)



L70 ANSWER 4 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2004:995903 HCAPLUS <>LOGINID: 20061121>>
 DOCUMENT NUMBER: 141:410698
 TITLE: Preparation of α -oxoacid-substituted phenols for
 the treatment of metabolic disorders
 INVENTOR(S): Hodge, Kirvin L.; Sharma, Shalini;
 Von Borstel, Reid W.
 PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA; Von Borstel,
 Reid W.
 SOURCE: PCT Int. Appl., 52 pp.
 CODEN: PIIXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004098496	A2	20041118	WO 2004-US12141	20040420
WO 2004098496	A3	20050331		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2004237602	A1	20041118	AU 2004-237602	20040420
CA 2522738	AA	20041118	CA 2004-2522738	20040420
EP 1617835	A2	20060125	EP 2004-750363	20040420
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
CN 1780614	A	20060531	CN 2004-80011552	20040420
JP 2006525331	T2	20061109	JP 2006-513150	20040420
PRIORITY APPLN. INFO.:			US 2003-466663P	P 20030430
			WO 2004-US12141	W 20040420
OTHER SOURCE(S): GI	CASREACT 141:410698; MARPAT 141:410698			



AB Title compds. I [n = 1-2; m = 0-4; q, p = 0-1; R₂ = alkyl; R₃ = H, halo; A = (un)substituted Ph, cycloalkyl, etc.; R₁ = H, alkyl] are prepared For instance, 2-oxo-2-[3-(2,6-dimethylbenzyloxy)phenyl]acetic acid (II) is prepared by SeO₂ oxidation of the corresponding ethanone precursor (prior art). II showed a statistically significant decrease in blood glucose and triglycerides in obese mice compared to control at 60 mg/Kg. I are useful for the treatment of various metabolic disorders, such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis.

IC ICM A61K

CC 25-10 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
Section cross-reference(s): 1, 63

L70 ANSWER 5 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:927013 HCPLUS <>LOGINID: 20061121>>

DOCUMENT NUMBER: 141:395291

TITLE: Preparation of benzyloxyphenyl acids and related compounds for the treatment of metabolic disorders

INVENTOR(S): *Hodge, Kirvin L.; Kaufman, Robert J.; Lee, Albert; Sharma, Shalini; Von Boistel, Reid W.*

PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA

SOURCE: PCT Int. Appl., 42 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

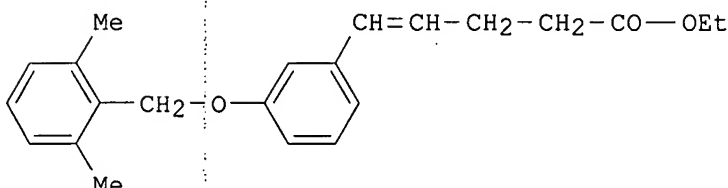
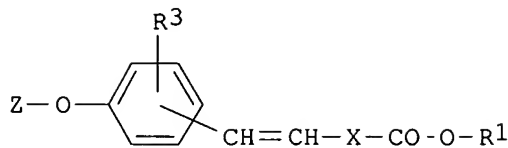
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004093806	A2	20041104	WO 2004-US12142	20040420
WO 2004093806	A3	20050407		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2521589	AA	20041104	CA 2004-2521589	20040420
EP 1618086	A2	20060125	EP 2004-750364	20040420
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR				
CN 1777576	A	20060524	CN 2004-80010732	20040420
JP 2006524252	T2	20061026	JP 2006-513151	20040420
PRIORITY APPN. INFO.:			US 2003-464553P	P 20030422
			WO 2004-US12142	W 20040420

OTHER SOURCE(S): MARPAT 141:395291

GI



AB Title compds. I [Z = $(\text{CH}_2)_n(\text{NR}^3)q(\text{CH}_2)^t\text{A}$; X = $(\text{CH}_2)_m$; R¹ = H, alkyl; R² = alkyl; R³ = H, halo, alkyl, etc.; n = 1-2; m = 2-3; q = 0-1; t = 0-1; A = (un)substituted Ph, cycloalkyl, heteroarom., etc.] and their pharmaceutically acceptable salts were prepared. For example, condensation of 3-(2,6-dimethylbenzyloxy)benzaldehyde and triphenylethylbutyrate phosphonium bromide afforded claimed benzyloxyphenyl acid ester II in 62% yield. In serum glucose assays in b/db mice, compound II exhibited glucose mg/dL of 651 at 100 mg/kg dosage. Compds. I are claimed useful for the treatment of metabolic disorders, i.e., diabetes, metabolic syndrome X, obesity, etc.

IC ICM A61K

CC 25-18 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
Section cross-reference(s): 1

L70 ANSWER 6 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:902090 HCAPLUS <>LOGINID::20061121>>

DOCUMENT NUMBER: 141:384282

TITLE: Compounds for the treatment of metabolic disorders

INVENTOR(S): *Hodge, Kirvin L.; Sharma, Shalini;
Von Borstel, Reid W.; Wolpe, Stephen
D.*

PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA

SOURCE: PCT Int. Appl., 47 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004091486	A2	20041028	WO 2004-US10799	20040408
WO 2004091486	A3	20050120		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW			
RW:	BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,			

BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2004229418	A1	20041028	AU 2004-229418	20040408
CA 2521621	AA	20041028	CA 2004-2521621	20040408
EP 1633340	A2	20060315	EP 2004-759257	20040408
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK				
BR 2004009469	A	20060418	BR 2004-9469	20040408
CN 1774244	A	20060517	CN 2004-80010105	20040408
JP 2006523696	T2	20061019	JP 2006-509802	20040408
US 2006014784	A1	20060119	US 2005-531618	20050414
NO 2005004791	A	20051220	NO 2005-4791	20051018
PRIORITY APPLN. INFO.:			US 2003-462960P	P 20030415
			WO 2004-US10799	W 20040408

OTHER SOURCE(S): MARPAT 141:384282

AB Agents such as 4-(3-(2,6-dimethylbenzyloxy)phenyl)-4-hydroxybutanoic acid, useful for the treatment of various metabolic disorders, such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis are disclosed. Thus, 4-(3-(2,6-dimethylbenzyloxy)phenyl)-4-(R)-hydroxybutanoic acid was prepared by the NaBH4 reduction of 4-(3-(2,6-dimethylbenzyloxy)phenyl)-4-oxobutanoic acid. The above compound elicited a significant reduction in blood glucose.

IC ICM A61K

CC 63-6 (Pharmaceuticals)

L70 ANSWER 7 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:718293 HCPLUS <>LOGINID::20061121>>

DOCUMENT NUMBER: 141:236676

TITLE: Compounds for the treatment of metabolic disorders

INVENTOR(S): *Hodge, Kirvin L.; Lee, Albert; Sharma, Shalini; Von Borstel, Reid W.*

PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA

SOURCE: PCT Int. Appl., 78 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004073611	A2	20040902	WO 2004-US3718	20040209
WO 2004073611	A3	20041125		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2004212905	A1	20040902	AU 2004-212905	20040209
CA 2513092	AA	20040902	CA 2004-2513092	20040209
EP 1601251	A2	20051207	EP 2004-709467	20040209
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK				

BR 2004007506	A	20060214	BR 2004-7506	20040209
CN 1750758	A	20060322	CN 2004-80004150	20040209
JP 2006517920	T2	20060803	JP 2005-518490	20040209
US 2006247309	A1	20061102	US 2005-531630	20050414
NO 2005003211	A	20051020	NO 2005-3211	20050630
PRIORITY APPLN. INFO.:			US 2003-447168P	P 20030213
			WO 2004-US3718	W 20040209

OTHER SOURCE(S): MARPAT 141:236676

AB Agents useful for the treatment of various metabolic disorders, such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis are disclosed. Formula (I) wherein n is 1 or 2; m is 0, 1, 2, 4 or 5; q is 0 or 1; t is 0 or 1; R₂ is alkyl from 1 to 3 carbon atoms; R₃ is hydrogen, halo, alkyl having from 1 to 3 carbon atoms, or alkoxy having from 1 to 3 carbon atoms; A is Ph, unsubstituted or substituted by one or two groups selected from: halo, alkyl having 1 or 2 carbon atoms, perfluoromethyl, alkoxy having 1 or 2 carbon atoms, and perfluoromethoxy; or cycloalkyl having from 3 to 6 ring carbon atoms wherein the cycloalkyl is unsubstituted or one or two ring carbons are independently mono-substituted by Me or ethyl; or a 5 or 6 membered heteroarom. ring having 1 or 2 ring heteroatoms selected from N, S and O and the heteroarom. ring is covalently bound to the remainder of the compds. of formula (I) by a ring carbon; and R₁ is hydrogen or alkyl having 1 or 2 carbon atoms. Alternatively, when R₁ is hydrogen, the biol. active agent can be a pharmaceutically acceptable salt of the compound of Formula (I).

IC ICM A61K

CC 1-10 (Pharmacology)

Section cross-reference(s): 25

L70 ANSWER 8 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:696360 HCAPLUS <>LOGINID::20061121>>

DOCUMENT NUMBER: 141:225492

TITLE: Preparation of isoxazoles as inhibitors of heat shock proteins

INVENTOR(S): Drysdale, Martin James; Dymock, Brian William; Finch, Harry; Webb, Paul; McDonald, Edward; James, Karen Elizabeth; Cheung, Kwai Ming; Mathews, Thomas Peter

PATENT ASSIGNEE(S): Vernalis Cambridge Limited, UK; Cancer Research Technology Ltd; The Institute of Cancer Research; et al.; et al.

SOURCE: PCT Int. Appl., 180 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

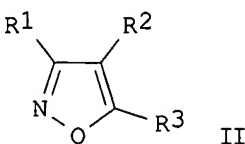
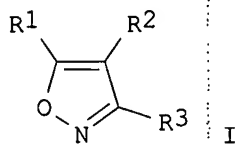
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2004072051	A1	20040826	WO 2004-GB506	20040209
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI				
RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2004210779	A1	20040826	AU 2004-210779	20040209

CA 2515726	AA 20040826	CA 2004-2515726	20040209
EP 1611112	A1 20060104	EP 2004-709273	20040209
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK			
CN 1771235	A 20060510	CN 2004-80009339	20040209
JP 2006517572	T2 20060727	JP 2006-502254	20040209
BR 2004007403	A 20061003	BR 2004-7403	20040209
NO 2005004195	A 20051109	NO 2005-4195	20050909
US 2006241106	A1 20061026	US 2006-544443	20060421
PRIORITY APPLN. INFO.:		GB 2003-3105	A 20030211
		GB 2003-6560	A 20030321
		GB 2003-13751	A 20030613
		WO 2004-GB506	W 20040209

OTHER SOURCE(S): MARPAT 141:225492
GI



AB Title compds. [I, II; R1 = Ar1(Alk1)p(Z)r(Alk2)sQ; Ar1 = (substituted) aryl, heteroaryl; Alk1, Alk2 = (substituted) alkylene, alkenylene; p, r, s = 0, 1; Z = O, S, CO, CS, SO2, CO2, CONRA, CSNRA, SO2NRA, NRACO, NRASO2, NRA; RA = H, alkyl; Q = H, (substituted) carbocyclyl, heterocyclyl; R2 = Ar1(Alk1)p(Z)r(Alk2)sQ, carboxamide, carbocyclyl, heterocyclyl optionally substituted by (Alk1)pZr(Alk2)sQ; R3 = H, (substituted) cycloalkyl, cycloalkenyl, alkyl, alkenyl, alkynyl, carboxyl, carboxamide, carboxyl ester], were prepared. Thus, NH2OH.HCl and 7-hydroxy-3-(4-methoxyphenyl)-2-methylchromen-4-one (preparation given) were refluxed 4 h in pyridine to give 4-[4-(4-methoxyphenyl)-3-methylisoxazol-5-yl]benzene-1,3-diol. The latter in the Malachite Green ATPase assay inhibited HSP90 with IC50 <50 µM.

IC ICM C07D261-08
ICS C07D413-04; C07D413-10; C07D417-04; C07D261-10; C07D495-04;
A61P035-00

CC 28-6 (Heterocyclic Compounds (More Than One Hetero Atom))
Section cross-reference(s): 1, 63

IT **Eye, disease**
(diabetic **retinopathy**, treatment; preparation of isoxazoles as inhibitors of heat shock proteins)

IT Autoimmune disease
(insulin-dependent **diabetes mellitus**, treatment; preparation of isoxazoles as inhibitors of heat shock proteins)

IT **Diabetes mellitus**
(insulin-dependent, treatment; preparation of isoxazoles as inhibitors of heat shock proteins)

IT 487-49-0P	2284-30-2P	13004-42-7P	19337-03-2P	22877-01-6P
23504-03-2P	29048-54-2P	90110-32-0P	103620-87-7P	130307-08-3P
140660-31-7P	328018-52-6P	536974-86-4P	558645-35-5P	705963-54-8P
747412-81-3P	747412-82-4P	747412-94-8P	747413-00-9P	
747413-03-2P	747413-04-3P	747413-05-4P	747413-06-5P	
747413-07-6P	747413-12-3P	747413-16-7P	747413-17-8P	747413-18-9P
747413-19-0P	747413-21-4P	747413-22-5P	747413-23-6P	747413-31-6P
747413-33-8P	747413-34-9P	747413-35-0P	747413-68-9P	

747413-69-0P 747413-70-3P 747413-71-4P 747413-72-5P
747413-73-6P 747413-74-7P 747413-75-8P 747414-06-8P 747414-07-9P
747414-08-0P 747414-09-1P 747414-10-4P 747414-11-5P 747414-12-6P
747414-16-0P 747414-17-1P 747414-18-2P 747414-19-3P 747414-20-6P
747414-21-7P 747414-22-8P 747414-23-9P 747414-24-0P 747414-48-8P
747414-49-9P 747414-50-2P 747414-51-3P 747414-52-4P 747414-53-5P
747414-55-7P 747414-56-8P 747414-57-9P 747414-62-6P 747414-63-7P
747414-64-8P 747414-65-9P 747414-66-0P 747414-67-1P 747414-69-3P
747414-70-6P 747414-71-7P 747414-72-8P 747414-73-9P
747414-74-0P 747414-76-2P 747414-78-4P 747414-79-5P 747414-80-8P
747414-84-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of isoxazoles as inhibitors of heat shock proteins)

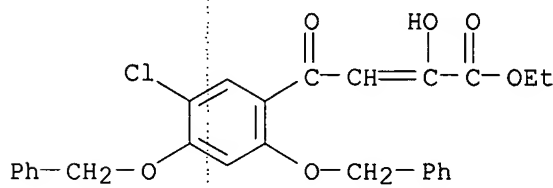
IT 747413-03-2P 747413-69-0P 747414-70-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of isoxazoles as inhibitors of heat shock proteins)

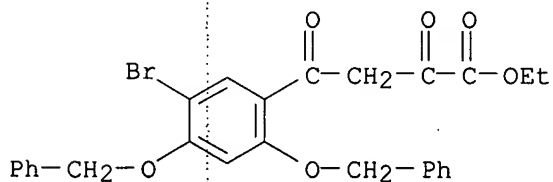
RN 747413-03-2 HCPLUS

CN 2-Butenoic acid, 4-[5-chloro-2,4-bis(phenylmethoxy)phenyl]-2-hydroxy-4-oxo-, ethyl ester (9CI) (CA INDEX NAME)



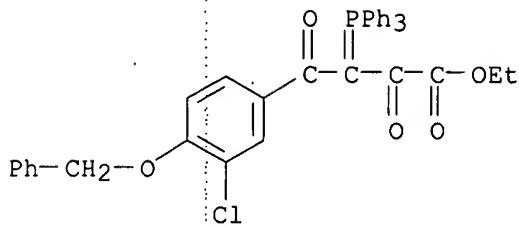
RN 747413-69-0 HCPLUS

CN Benzenebutanoic acid, 5-bromo- α,γ -dioxo-2,4-bis(phenylmethoxy)-, ethyl ester (9CI) (CA INDEX NAME)



RN 747414-70-6 HCPLUS

CN Benzenebutanoic acid, 3-chloro- α,γ -dioxo-4-(phenylmethoxy)-β-(triphenylphosphoranylidene)-, ethyl ester (9CI) (CA INDEX NAME)



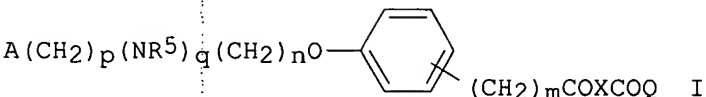
L70 ANSWER 9 OF 15 HCAPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 2002:964135 HCAPLUS <>LOGINID::20061121>>
 DOCUMENT NUMBER: 138:24543
 TITLE: Preparation of benzyloxyphenyloxobutyrates and related compounds for the treatment of metabolic disorders
 INVENTOR(S): *Sharma, Shalini; Von Borstel, Reid W.; Hodge, Kirvin L.*
 PATENT ASSIGNEE(S): Wellstat Therapeutics Corporation, USA; Bamat, Michael K.
 SOURCE: PCT Int. Appl., 242 pp.
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002100341	A2	20021219	WO 2002-US18388	20020612
WO 2002100341	A3	20040701		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 2450221	AA	20021219	CA 2002-2450221	20020612
US 2003149107	A1	20030807	US 2002-167839	20020612
US 7101910	B2	20060905		
EP 1461323	A2	20040929	EP 2002-744271	20020612
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR				
JP 2005501012	T2	20050113	JP 2003-503168	20020612
CN 1608055	A	20050420	CN 2002-811881	20020612
BR 2002010383	A	20060404	BR 2002-10383	20020612
US 2004077896	A1	20040422	US 2003-684644	20031014
US 6924314	B2	20050802		
US 2004092518	A1	20040513	US 2003-684735	20031014
US 7041659	B2	20060509		
US 2004092516	A1	20040513	US 2003-685183	20031014
US 6946491	B2	20050920		
US 2004097585	A1	20040520	US 2003-684730	20031014
US 6916848	B2	20050712		

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US 2004236100	A1	20041125	US 2003-684660	20031014
US 6858602	B2	20050222		
US 2004267025	A1	20041230	US 2003-684740	20031014
US 7045541	B2	20060516		
ZA 2003009627	A	20050617	ZA 2003-9627	20031211
US 2004242692	A1	20041202	US 2004-865088	20040610
US 2005004115	A1	20050106	US 2004-892950	20040716
US 7012071	B2	20060314		
US 2005090555	A1	20050428	US 2004-5449	20041206
US 2005256333	A1	20051117	US 2005-481042	20050114
PRIORITY APPLN. INFO.:				
			US 2001-297282P	P 20010612
			US 2002-167839	A3 20020612
			WO 2002-US18388	W 20020612
			US 2003-685183	A3 20031014
			US 2004-865088	A1 20040610

OTHER SOURCE(S): MARPAT 138:24543
GI



AB Biol. active title compds. [I; n = 1, 2; m, q, p = 0, 1; R⁵ = alkyl; R⁹ = H, halo, alkoxy; A = (halo-, alkyl-, perfluoromethyl-, alkoxy-, perfluoromethoxy-substituted) Ph, (Me-, Et-substituted) cycloalkyl, 5-6 membered heteroarom. ring having 1-2 N, S, O atoms; X = CH₂, Q = OR₁, R₁ = Et; or X = CH₂CR₁R₁₃, CH₂CH(NHAc), Q = OR₁, R₁ = H, alkyl; or X = CH₂CH₂, Q = NR₁₀R₁₁; R₁₂, R₁₃ = H, Me; 1 of R₁₀, R₁₁ = H, alkyl, OH, the other = H, alkyl], were prepared Thus, 4-(2-fluorobenzyl)acetophenone (preparation given) in THF and DMPU was treated with a solution of Li bis(trimethylsilyl)amide at -60°; after 10 min, tert-Bu bromoacetate was added followed by stirring for an addnl. 10 min and warming to room temperature for 4 h to give tert-Bu[4-[4-(2-fluorobenzyl)phenyl]-4-oxobutyrate. The latter was stirred with CF₃CO₂H in CH₂Cl₂ to give 4-[4-(2-fluorobenzyl)phenyl]-4-oxobutyric acid. Tested I showed antidiabetic activity in a variety of tests. I are useful in treatment of various metabolic disorders such as insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis and arteriosclerosis.

IC ICM A61K

CC 25-17 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
Section cross-reference(s): 1, 27, 28

L70 ANSWER 10 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:676588 HCPLUS <<LOGINID:20061121>>
DOCUMENT NUMBER: 135:221312
TITLE: Therapeutic uses of PPAR mediators as ABC-1 expression modulators, and preparation thereof
INVENTOR(S): Jaye, Michael; Duverger, Nicolas; Searfoss, George; Minnich, Anne
PATENT ASSIGNEE(S): Aventis Pharma Deutschland G.m.b.H., Germany
SOURCE: PCT Int. Appl., 176 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent

Zhang 10/532690

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001066098	A2	20010913	WO 2001-EP2482	20010306
WO 2001066098	A3	20020404		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
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BR 2001009107	A	20021203	BR 2001-9107	20010306
EP 1267874	A2	20030102	EP 2001-956185	20010306
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ZA 2002007061	A	20031114	ZA 2002-7061	20020903
NO 2002004273	A	20021007	NO 2002-4273	20020906
US 2003220373	A1	20031127	US 2002-237578	20020909
PRIORITY APPLN. INFO.:			US 2000-188323P	P 20000309
			GB 2000-13589	A 20000606
			WO 2001-EP2482	W 20010306

OTHER SOURCE(S): MARPAT 135:221312

AB The invention discloses the use of PPAR mediators, and their pharmaceutical compns., as ATP binding cassette transporter 1 (ABC-1) expression modulators, wherein the PPAR ligand receptor agonists of the invention are useful as inducers of ABC-1 expression. Preparation of compds. of the invention is included. Also disclosed are methods for treating e.g. low levels of HDL.

IC ICM A61K031-00

CC 1-10 (Pharmacology)

Section cross-reference(s): 27, 28, 63

IT 120128-20-3P	123225-56-9P	123225-57-0P	123225-58-1P	123225-59-2P
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RL: **BAC** (*Biological activity or effector, except adverse*); BSU
(*Biological study, unclassified*); SPN (*Synthetic preparation*); **THU**
(*Therapeutic use*); BIOL (*Biological study*); PREP (*Preparation*); USES
(*Uses*)

(PPAR mediators as ABC-1 expression modulators, preparation, and therapeutic use)

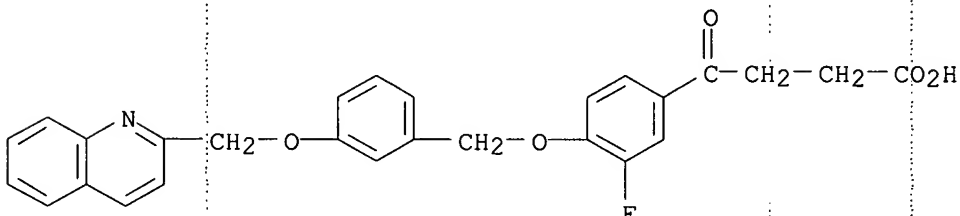
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(*Biological study, unclassified*); SPN (*Synthetic preparation*); **THU**
(*Therapeutic use*); BIOL (*Biological study*); PREP (*Preparation*); USES
(*Uses*)

(PPAR mediators as ABC-1 expression modulators, preparation, and therapeutic use)

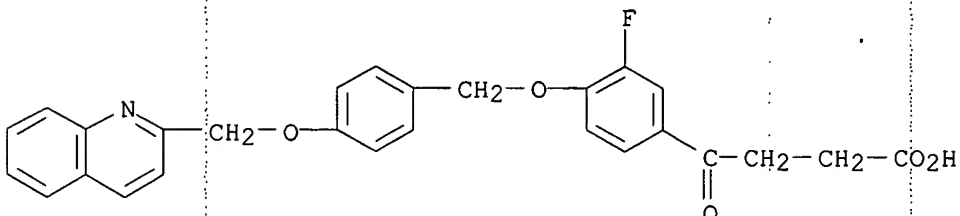
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CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[(3-(2-quinolinylmethoxy)phenyl)methoxy]- (9CI) (CA INDEX NAME)



RN 223772-46-1 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[(4-(2-quinolinylmethoxy)phenyl)methoxy]- (9CI) (CA INDEX NAME)



L70 ANSWER 11 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2000:772613 HCPLUS <<LOGINID: 20061121>>

DOCUMENT NUMBER: 133:335164

TITLE: Tri-aryl acid derivatives as PPAR receptor ligands

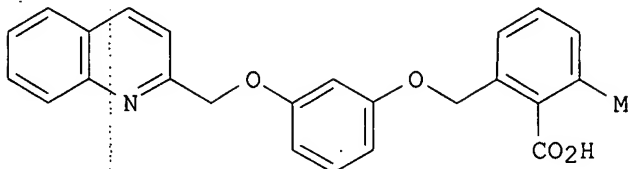
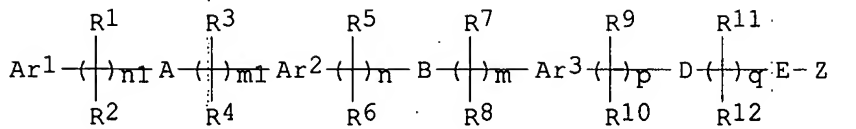
Zhang 10/532690

INVENTOR(S): Jayyosi, Zaid; McGeehan, Gerard M.; Kelley, Michael F.; Labaudiniere, Richard F.; Zhang, Litao; Caulfield, Thomas J.; Minnich, Anne; Bobko, Mark; Morris, Robert; Groneberg, Robert D.; McGarry, Daniel G.
PATENT ASSIGNEE(S): Aventis Pharmaceuticals Products Inc., USA
SOURCE: PCT Int. Appl., 257 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000064876	A1	20001102	WO 2000-US11490	20000428
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW				
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HR 2001000793	A1	20030228	HR 2001-793	20011026
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PRIORITY APPLN. INFO.:			US 1999-131454P	P 19990428
			WO 2000-US11490	W 20000428

OTHER SOURCE(S): MARPAT 133:335164

GI



AB This invention is directed to triaryl acid derivs. I and their salts, N-oxides, hydrates, solvates, and pharmaceutical compns. [wherein: Ar1, Ar2, Ar3 = aryl, fused arylcycloalkenyl, fused arylcycloalkyl, fused arylheterocyclenyl, fused arylheterocyclyl; heteroaryl, fused heteroarylcyclenyl, fused heteroarylcyclkemyl, fused heteroarylcyclalkyl, fused heteroarylheterocyclenyl, or fused heteroarylheterocyclyl; A = bond, O, S, SO, SO₂, CO, (un)substituted NH, NHCO, CONH, NHCONH, CH:N, etc.; B = bond, O, S, SO, SO₂, C.tplbond.C, CO, (un)substituted NH, NHCO, or CONH; D = bond, O, S, C.tplbond.C, CO, (un)substituted NH, NHCO, or CONH; E = bond, CH₂CH₂; Z = (un)substituted CO₂H, CHO, cyclo-imide, cyano, sulfonylaminocarbonyl, sulfonylamino, carbamoyl, tetrazolyl, etc.; R1, R3, R5, R7, R9, R11 = H, halo, alkyl, CO₂H, alkoxy carbonyl, aralkyl; R2, R4, R6, R8, R10, R12 = (CH₂)₀₋₃X (where X = H or various substituents); n1 = 0-4; m1 = 0-4; n = 0-4; m = 0-5; p = 0-4; q = 0-6; with numerous provisos]. The compds. are PPAR receptor ligands, useful as agonists or antagonists thereof (no data). For instance, 2,6-dimethylbenzoic acid underwent a sequence of: (1) Me esterification, (2) benzylic monobromination, (3) etherification with 3-(quinolin-2-ylmethoxy)phenol, and (4) alkaline hydrolysis with NaOH in aqueous EtOH, to give title compound

II.**ICM** C07D215-14**ICS** A61K031-33; A61K031-19; A61P043-00; C07D401-12; C07D401-14; C07D215-18; C07D405-12; C07D263-32; C07D213-30; C07D241-42; C07D277-24; C07D261-08; C07D271-06; C07D277-64**CC** 27-17 (Heterocyclic Compounds (One Hetero Atom))

Section cross-reference(s): 1

IT *Diabetes mellitus*

(non-insulin-dependent, treatment of; preparation of tri-aryl acid derivs. as PPAR receptor ligands)

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RL: **BAC** (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); **THU** (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of tri-aryl acid derivs. as PPAR receptor ligands)

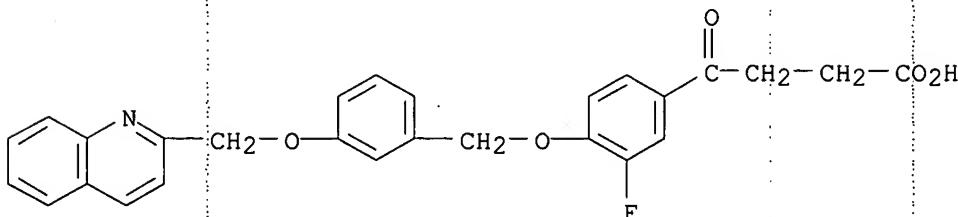
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RL: **BAC** (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); **THU** (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

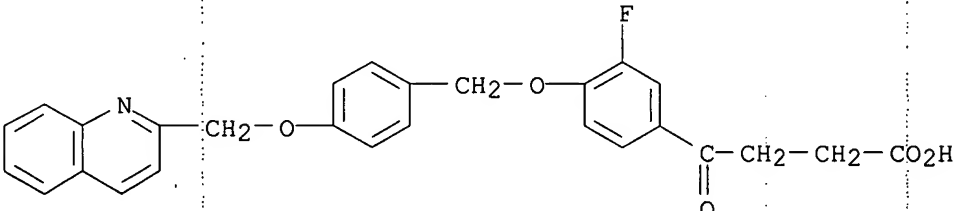
(preparation of tri-aryl acid derivs. as PPAR receptor ligands)

RN 223772-45-0 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[(3-(2-quinolinylmethoxy)phenyl)methoxy]- (9CI) (CA INDEX NAME)



RN 223772-46-1 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[[4-(2-quinolinylmethoxy)phenyl]methoxy]- (9CI) (CA INDEX NAME)

RETABLE

Referenced Author (RAU)	Year (R PY)	VOL (R VL)	PG (R PG)	Referenced Work (R WK)	Referenced File
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L70 ANSWER 12 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:282096 HCPLUS <>LOGINID::20061121>>

DOCUMENT NUMBER: 130:320864

TITLE: PPAR- γ -binding quinoline derivatives, their preparation, and their therapeutic use

INVENTOR(S): Jayyosi, Zaid; McGeehan, Gerard M.; Kelley, Michael F.

PATENT ASSIGNEE(S): Rhone-Poulenc Rorer Pharmaceuticals Inc., USA

SOURCE: PCT Int. Appl., 125 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 8

PATENT INFORMATION:

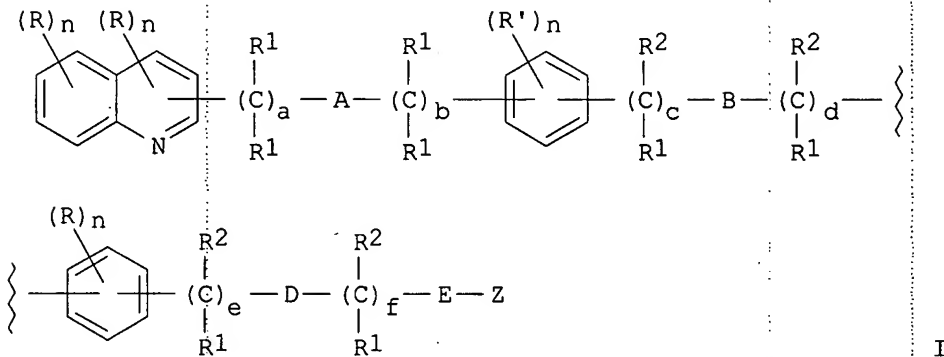
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9920275	A1	19990429	WO 1998-US21947	19981016
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OTHER SOURCE(S): MARPAT 130:320864

GI



AB A method for mediating the activity of PPAR- γ receptor comprises contacting the PPAR- γ receptor with I [A = O, S, (R1)C=C(R1), bond; B = O, S, SO, SO₂, NR1, bond; D = O, S, NR1, (R1)C=C(R1), bond; E = bond; a = 0-2; b = 0, 1; c = 0-4; d = 0-5; e = 0-4; f = 0-5; n = 0-2; R = H; R' = H; R1 = H; R2 = (CH₂)qX, or two vicinal R2 taken together with the carbon atoms through which the two vicinal R2 are linked form cycloalkylene, etc.; q = 0-3; X = H]. Preparation of I is described. The compds. may be used to treat cardiovascular conditions, **diabetes**, **hyperlipidemia**, **hypertension**, eating disorders, etc.

IC ICM A61K031-47

ICS A61K031-38; A61K031-35; A61K031-155; A61K031-18

CC 1-12 (Pharmacology)

Section cross-reference(s): 27, 28

IT 114497-47-1P	123225-56-9P	123225-57-0P	123225-58-1P	123225-59-2P
123225-60-5P	123225-61-6P	123225-63-8P	123225-64-9P	123225-69-4P
123225-71-8P	123225-72-9P	123225-76-3P	123225-82-1P	123225-94-5P
123225-95-6P	123225-96-7P	123225-98-9P	123225-99-0P	123226-00-6P
123226-01-7P	123226-03-9P	123226-04-0P	123226-05-1P	123226-06-2P
123226-07-3P	123226-08-4P	123226-09-5P	123226-10-8P	123226-11-9P
123226-13-1P	123226-14-2P	123226-15-3P	123226-16-4P	123226-17-5P

123226-19-7P 123226-20-0P 123226-21-1P 123226-25-5P 123226-27-7P
 123247-23-4P 123247-25-6P 123247-27-8P 123247-28-9P 123692-25-1P
 123692-29-5P 123692-37-5P 123692-38-6P 123692-39-7P 123791-11-7P
 123791-15-1P 124993-46-0P 124993-48-2P 128760-62-3P 128760-70-3P
 128760-73-6P 129649-40-7P 223772-08-5P 223772-12-1P 223772-14-3P
 223772-15-4P 223772-18-7P 223772-26-7P 223772-42-7P 223772-43-8P
 223772-44-9P 223772-45-0P 223772-46-1P 223772-47-2P
 223772-48-3P

RL: **BAC** (*Biological activity or effector, except adverse*); BSU
 (*Biological study, unclassified*); SPN (*Synthetic preparation*); THU
 (*Therapeutic use*); BIOL (*Biological study*); PREP (*Preparation*); USES
 (*Uses*)

(PPAR- γ -binding quinoline derivative preparation and therapeutic use)

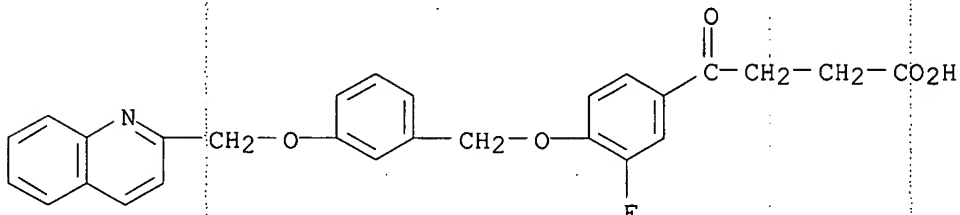
IT 223772-45-0P 223772-46-1P

RL: **BAC** (*Biological activity or effector, except adverse*); BSU
 (*Biological study, unclassified*); SPN (*Synthetic preparation*); THU
 (*Therapeutic use*); BIOL (*Biological study*); PREP (*Preparation*); USES
 (*Uses*)

(PPAR- γ -binding quinoline derivative preparation and therapeutic use)

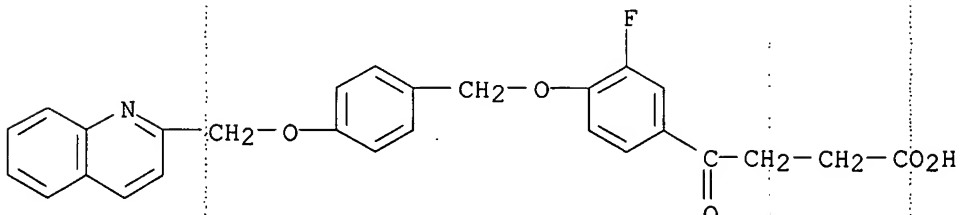
RN 223772-45-0 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[[3-(2-quinolinylmethoxy)phenyl]methoxy]- (9CI) (CA INDEX NAME)



RN 223772-46-1 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -oxo-4-[[4-(2-quinolinylmethoxy)phenyl]methoxy]- (9CI) (CA INDEX NAME)



RETABLE

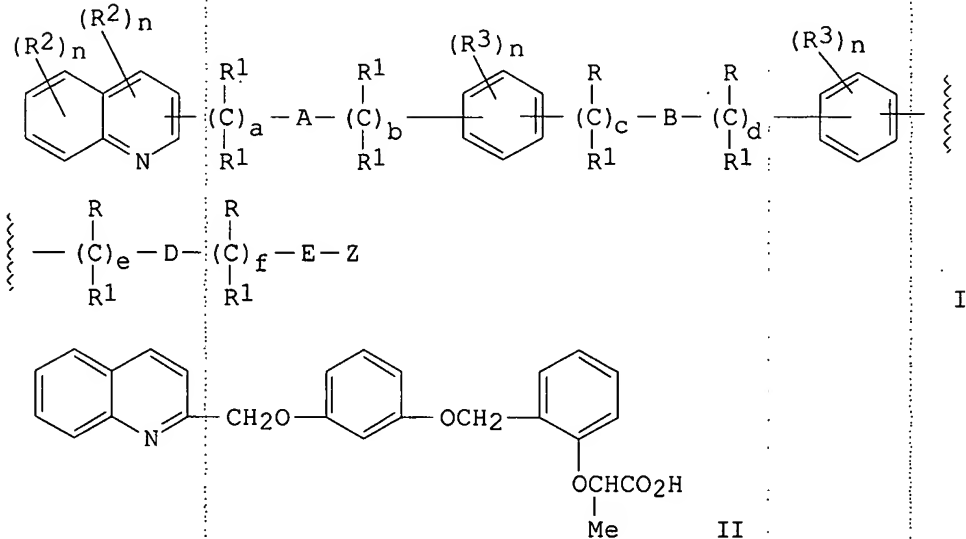
Referenced (RAU)	Author (RPY)	Year (RVL)	VOL (RPG)	PG (RWK)	Referenced Work (EP, WO, US)	Referenced File (HCPLUS)
Asahi Glass Company Ltd		1996			EP 0709377 A1	
Merrell Dow Pharmaceuti		1995			WO 9514669 A1	
Sterne		1965			US 3174901 A	

Zhang 10/532690

L70 ANSWER 13 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1990:497460 HCPLUS <<LOGINID: 20061121>>
DOCUMENT NUMBER: 113:97460
TITLE: Preparation of quinoline derivatives useful as
lipoxygenase inhibitors and/or leukotriene antagonists
INVENTOR(S): Huang, Fu Chi; Galembo, Robert Anthony, Jr.; Campbell,
Henry Flud
PATENT ASSIGNEE(S): Rorer International (Overseas), Inc., USA
SOURCE: Eur. Pat. Appl., 30 pp.
CODEN: EPXXDW
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 5
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 348155	A1	19891227	EP 1989-306232	19890620
EP 348155	B1	19990512		
R: DE, ES, FR, GB, IT				
US 4920131	A	19900424	US 1988-209428	19880621
EP 784052	A1	19970716	EP 1997-200638	19890620
EP 784052	B1	20040901		
R: DE, ES, FR, GB, IT				
US 5059610	A	19911022	US 1990-477896	19900420
PRIORITY APPLN. INFO.:			US 1988-209428	A 19880621
			US 1987-116420	A2 19871103
			US 1987-116428	A2 19871103
			US 1987-116597	A2 19871103
			WO 1988-US3897	W 19881101
			EP 1989-306232	A3 19890620

OTHER SOURCE(S): MARPAT 113:97460
GI



AB Title compds. I [A = O, S; B = bond, O, S, SO, SO₂, NR₁, CO, NR₁CO, CONR₁, CR₁:CR₁; D = O, S, NR₁, CR₁:CR₁, bond; E = bond, CR₁:CR₁; a = 0-2; b = 0-1; c = 0-4; d = 0-5, e = 0-4; f = 0-5; n = 0-2; R₂ = H, alkyl, OH, alkoxy, CO₂H, carbalkoxy, halo NO₂, haloalkyl, cyano, acyl; R₃ = H, OH, alkoxy, halo, etc.; R₁ = H, alkyl, aralkyl; R = (CH₂)_xX, O(CH₂)_xX, S(CH₂)_xX, NR₁(CH₂)_xX; x = 0-3; X = H, alkyl, alkenyl, aryl, alkoxy, amino, cyano, tetrazolyl, CO₂R, etc.; (R)₂ = (CH₂)_y with y = 1-4; RR₁ = (CH₂)_z with z = 2-5; (R₁)₂, RR₁ = CHR₁; Z = CO₂R₁, cyano, CONHSO₂R₄ with R₄ = H, alkyl, Ph, etc.; CON(R₁)₂, OR₁, (un)substituted tetrazolyl] were prepared as antiinflammatory and antiallergic agents (no data). Thus, condensation of o-cresol with MeCHBr Co₂Et and bromination of the product with NBS gave 2-(BrCH₂)C₆H₄ OCHMeCO₂Et, which underwent condensation with 3-(2-quinolinylmethoxy)phenol and basic hydrolysis to give quinoline derivative II. Several addnl. preps. and numerous I are given.

ICM C07D215-14

ICS C07D215-18; C07D401-12; A61K031-47

CC 27-17 (Heterocyclic Compounds (One Hetero Atom))

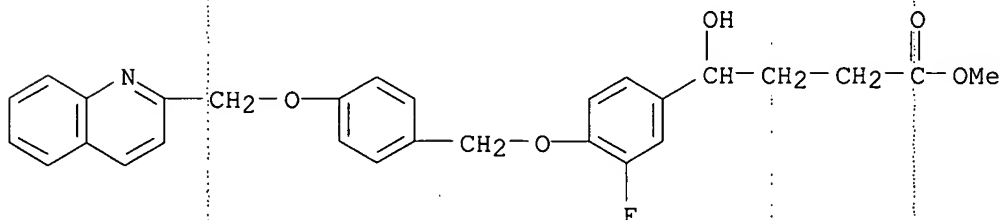
Section cross-reference(s): 1

IT	120128-20-3P	123247-27-8P	128760-39-4P	128760-40-7P	128760-41-8P
	128760-42-9P	128760-43-0P	128760-44-1P	128760-45-2P	128760-46-3P
	128760-47-4P	128760-48-5P	128760-49-6P	128760-50-9P	128760-51-0P
	128760-52-1P	128760-53-2P	128760-54-3P	128760-55-4P	128760-56-5P
	128760-57-6P	128760-58-7P	128760-59-8P	128760-60-1P	128760-61-2P
	128760-62-3P	128760-63-4P	128760-64-5P	128760-65-6P	128760-66-7P
	128760-67-8P	128760-68-9P	128760-69-0P	128760-70-3P	128760-71-4P
	128760-72-5P	128760-73-6P	128760-74-7P	128760-75-8P	128760-76-9P
	128760-86-1P	128760-87-2P	128784-91-8P	128784-92-9P	

128805-38-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of, as antiallergic and antiinflammatory agent)IT **128784-91-8P**RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of, as antiallergic and antiinflammatory agent)

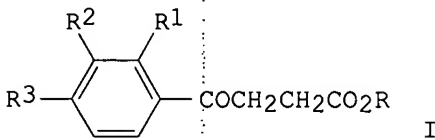
RN 128784-91-8 HCPLUS

CN Benzenebutanoic acid, 3-fluoro- γ -hydroxy-4-[(4-(2-quinolinylmethoxy)phenyl)methoxy]-, methyl ester (9CI) (CA INDEX NAME)

→ L70 ANSWER 14 OF 15 HCPLUS COPYRIGHT 2006 ACS on STN
 ACCESSION NUMBER: 1980:471315 HCPLUS <<LOGINID: 20061121>>
 DOCUMENT NUMBER: 93:71315
 TITLE: 2-(Substituted benzoyl)propionic acids
 INVENTOR(S): Noda, Kanji; Nakagawa, Akira; Hirano, Munehiko; Tsuji, Masayoshi; Ide, Hiroyuki
 PATENT ASSIGNEE(S): Hisamitsu Pharmaceutical Co., Inc., Japan
 SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF
 DOCUMENT TYPE: Patent
 LANGUAGE: Japanese
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 55015460	A2	19800202	JP 1978-89591	19780720
PRIORITY APPLN. INFO.: GI			JP 1978-89591	A 19780720



AB Seven I (R = H; R1 = Me2CH, H; R2 = H, NO2, Cl; R3 = Me2CH, EtO, cyclopropylmethoxy, allyloxy, etc.), having central-depressant, antiinflammatory, PCA-inhibitory and immunosuppressant activities (no data), were prepared by reacting succinic anhydride (II) with C6H6 derivs., or by reacting I (R = Et, R3 = OH) with R3Br (R3 = cyclopropylmethyl), followed by hydrolysis. Thus, m-(Me2CH)2C6H4 16.2 was added dropwise to II 10 and AlCl3 26.7 g in ClCH2CH2Cl with cooling, and the mixture was stirred 10 h at room temperature to give I (R = R2 = H, R1 = R3 = CHMe2).

IC C07C059-76; C07C079-46; C07C101-44

CC 25-18 (Noncondensed Aromatic Compounds)

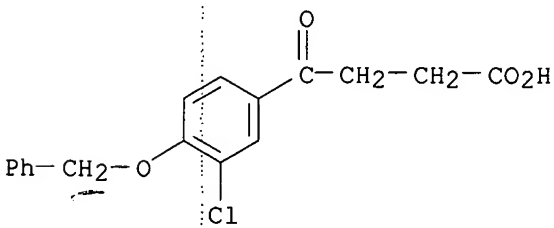
IT 74362-69-9P 74362-70-2P 74362-71-3P 74362-72-4P **74362-73-5P**
74362-74-6P 74391-08-5P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

IT **74362-73-5P**
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 74362-73-5 HCPLUS

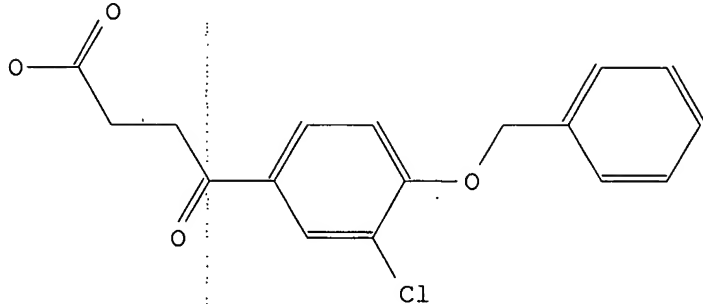
CN Benzenebutanoic acid, 3-chloro- γ -oxo-4-(phenylmethoxy)- (9CI) (CA INDEX NAME)



Zhang 10/532690

L53 ANSWER 1 OF 1 WPIX COPYRIGHT 2006
AN.S DCR-905973
DCSE 905973-0-0-0
CN.S 4-(4-Benzyl-3-chloro-phenyl)-4-oxo-butyric acid

THE THOMSON CORP on STN



MF C17 H15 Cl O4
SMF C17 H15 Cl O4 *1; TOTAL *1; TYPE *1
MW 318.7597
SDCN RAECKI

=> d all abeq tech 169 tot

L69 ANSWER 1 OF 1 WPIX COPYRIGHT 2006 THE THOMSON CORP on STN
AN 2004-411395 [38] WPIX <<LOGINID::20061121>>
DNC C2004-154379 [38]
TI Use of oxoalkanoates in the manufacture of a medicament for treatment of metabolic disorders e.g. insulin resistance syndrome and diabetes
DC B05
IN HODGE K L; SHARMA S; VON BORSTEL R W; WOLPE S D
PA (WELL-N) WELLSTAT THERAPEUTICS CORP; (HODG-I) HODGE K L; (SHAR-I) SHARMA S; (VBOR-I) VON BORSTEL R W; (WOLP-I) WOLPE S D
CYC 105
PI WO 2004041165 A2 20040521 (200438)* EN 22[0] A61K000-00
AU 2003286728 A1 20040607 (200469) EN A61K047-00
EP 1556085 A2 20050727 (200549) EN
US 20060035970 A1 20060216 (200614) EN
JP 2006507303 W 20060302 (200621) JA 19
AU 2003286728 A8 20051110 (200634) EN A61K047-00
ADT WO 2004041165 A2 WO 2003-US34185 20031028; US 20060035970 A1 Provisional
US 2002-423253P 20021101; AU 2003286728 A1 AU 2003-286728 20031028; EP
1556085 A2 EP 2003-777939 20031028; EP 1556085 A2 WO 2003-US34185
20031028; US 20060035970 A1 WO 2003-US34185 20031028; JP 2006507303 W WO
2003-US34185 20031028; JP 2006507303 W JP 2004-550151 20031028; US
20060035970 A1 US 2005-532690 20050426; AU 2003286728 A8 AU 2003-286728
20031028
FDT AU 2003286728 A1 Based on WO 2004041165 A; EP 1556085 A2 Based on
WO 2004041165 A; JP 2006507303 W Based on WO 2004041165 A; AU
2003286728 A8 Based on WO 2004041165 A
PRAI US 2002-423253P 20021101

US 2005-532690 20050426

IC ICM A61K005-; A61K047-00

IPCI A61K0031-185 [I,C]; A61K0031-192 [I,A]; A61K0031-21 [I,C]; A61K0031-235 [I,A]; A61K0031-185 [I,C]; A61K0031-192 [I,A]; A61K0031-21 [I,C]; A61K0031-216 [I,A]; A61P0001-00 [I,C]; A61P0001-16 [I,A]; A61P0013-00 [I,C]; A61P0013-12 [I,A]; A61P0017-00 [I,C]; A61P0017-02 [I,A]; A61P0025-00 [I,A]; A61P0027-00 [I,C]; A61P0027-02 [I,A]; A61P0027-12 [I,A]; A61P0003-00 [I,C]; A61P0003-04 [I,A]; A61P0003-06 [I,A]; A61P0003-10 [I,A]; A61P0009-00 [I,C]; A61P0009-10 [I,A]; A61P0009-12 [I,A]

AB WO 2004041165 A2 UPAB: 20060121

NOVELTY - Treatment of metabolic disorders involves administration of an agent.

DETAILED DESCRIPTION - Treatment of metabolic disorders involves administration of an agent selected from:

- (1) 4-(4-benzyloxy-3-chlorophenyl)-4-oxobutanoic acid;
- (2) methyl 4-(4-benzyloxy-2-methoxyphenyl)-4-oxobutanoate;
- (3) ethyl 4-(4-cyclohexylmethoxyphenyl)-4-oxobutanoate;
- (4) 4-(3-chloro-4-cyclopropylmethoxyphenyl)-4-oxobutanoic acid;
- (5) ethyl 3-(4-benzyloxyphenyl)-3-oxopropanoate;
- (6) ethyl 3-(3-benzyloxyphenyl)-3-oxopropanoate;
- (7) ethyl 3-(2-benzyloxyphenyl)-3-oxopropanoate;
- (8) methyl 3-(3-(2,6-dichlorobenzyloxy)phenyl)-3-oxopropanoate;
- (9) ethyl 3-(4-(4-chlorobenzyloxy)phenyl)-3-oxopropanoate;
- (10) ethyl 3-(3-(4-methoxybenzyloxy)phenyl)-3-oxopropanoate;
- (11) ethyl 3-(2-(4-methoxybenzyloxy)phenyl)-3-oxopropanoate;
- (12) ethyl 3-(2-(2-methoxybenzyloxy)phenyl)-3-oxopropanoate;
- (13) ethyl 3-(2-(3-methoxybenzyloxy)phenyl)-3-oxopropanoate;
- (14) ethyl 3-(4-benzyloxy-3-chlorophenyl)-3-oxopropanoate;
- (15) ethyl 3-(4-benzyloxy-3-methoxyphenyl)-3-oxopropanoate; or
- (16) ethyl 3-(3-benzyloxy-4-methoxyphenyl)-3-oxopropanoate.

ACTIVITY - Antidiabetic; Antiarteriosclerotic; Anorectic; Hypotensive; Antilipemic; Nephrotropic; Neuroprotective; Ophthalmological; Antiulcer; Immunomodulator.

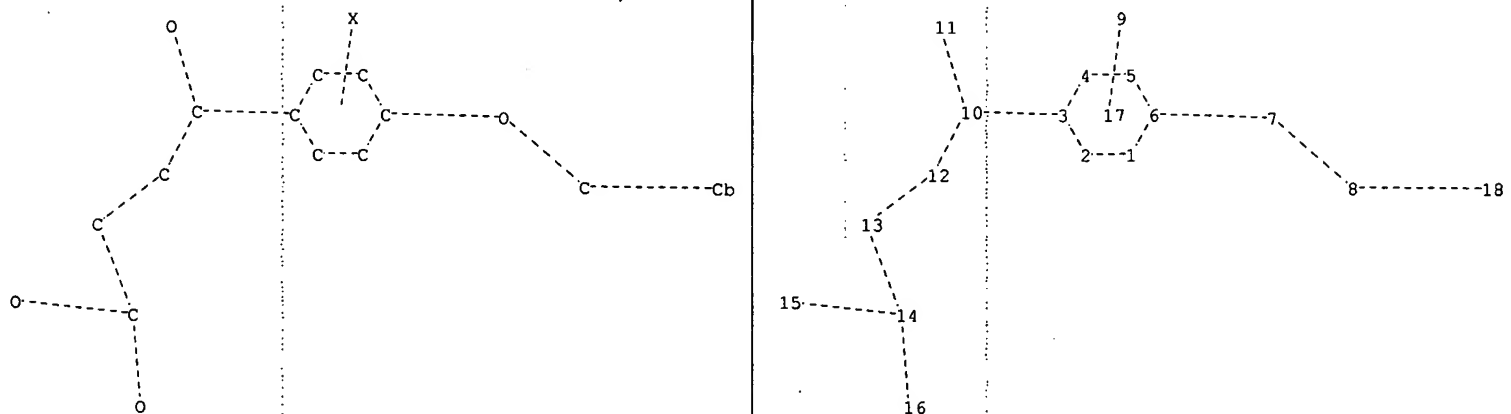
Test details described but no results given.

MECHANISM OF ACTION - None given.

USE - For treatment of metabolic disorders in a subject (such as human) e.g. insulin resistance syndrome and diabetes including Type I Diabetes and Type II Diabetes; for the treatment or reduction in the development of atherosclerosis, arteriosclerosis, obesity, hypertension, hyperlipidemia, fatty liver disease, nephropathy, neuropathy, retinopathy, foot ulceration and cataracts associated with diabetes; and for the treatment of cachexia (claimed).

ADVANTAGE - The agent effectively addresses the primary defects of insulin resistance and islet failure with fewer or milder side effects than existing drugs.

MC CPI: B10-C03; B10-F02; B14-D01E; B14-E11; B14-E12; B14-F02B; B14-F06; B14-F07; B14-J01; B14-N03; B14-N10; B14-N12; B14-N17B; B14-S04



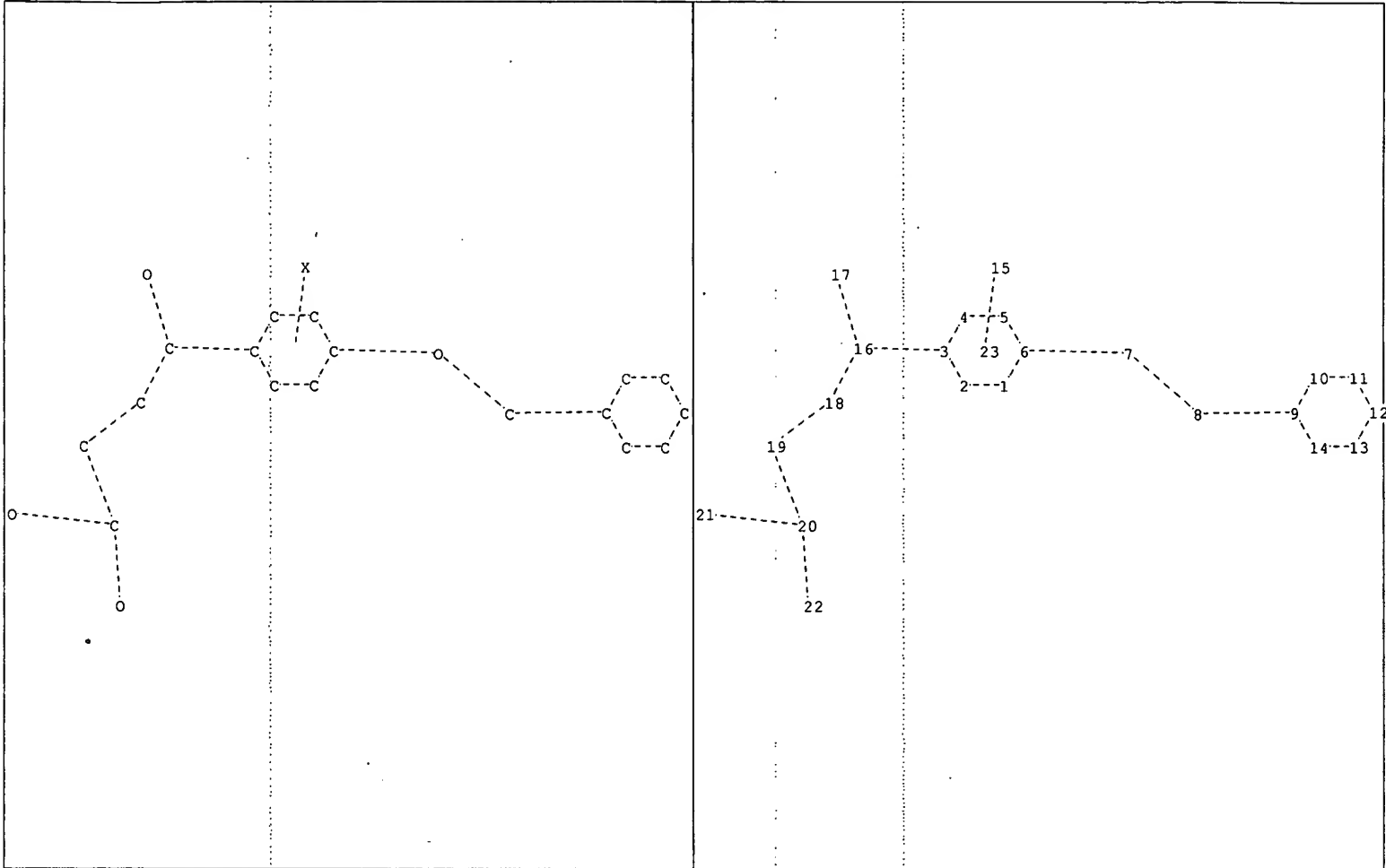
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ring nodes :
    1   2   3   4   5   6
chain bonds :
    3-10  6-7  7-8   8-18  10-11  10-12  12-13  13-14  14-15  14-16
ring bonds :
    1-2   1-6  2-3   3-4   4-5   5-6
exact/norm bonds :
    1-2   1-6  2-3   3-4   3-10  4-5   5-6   6-7   7-8   8-18  10-11  10-12  12-13
    13-14  14-15  14-16

Match level :
    1:Atom  2:Atom  3:Atom  4:Atom  5:Atom  6:Atom  7:CLASS  8:CLASS  9:CLASS
    10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS
    17:Atom  18:Atom
Element Count :
    Node 18: Limited
        C,C6

```

2nd Structure Search



chain nodes :

7 8 15 16 17 18 19 20 21 22

ring nodes :

1 2 3 4 5 6 9 10 11 12 13 14

chain bonds :

3-16 6-7 7-8 8-9 16-17 16-18 18-19 19-20 20-21 20-22

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 9-10 9-14 10-11 11-12 12-13 13-14

exact/norm bonds :

1-2 1-6 2-3 3-4 3-16 4-5 5-6 6-7 7-8 8-9 9-10 9-14 10-11 11-12
12-13 13-14 16-17 16-18 18-19 19-20 20-21 20-22

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:Atom
10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS 16:CLASS 17:CLASS
18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:CLASS 23:Atom

lst
The structure I searched

Day : Tuesday
 Date: 11/21/2006

Time: 10:46:16

PALM INTRANET**Inventor Information for 10/532690**

Inventor Name	City	State/Country
HODGE, KIRVIN L. 141	LAUREL	MARYLAND
SHARMA, SHALINI 142	GAITHERSBURG	MARYLAND
VON BORSTEL, REID W. 143	POTOMAC	MARYLAND
WOLPE, STEPHEN D. 146	BOYDS	MARYLAND

[Appln Info](#)[Contents](#)[Petition Info](#)[Atty/Agent Info](#)[Continuity/Reexam](#)[Foreign](#)

Search Another: Application# or Patent#
 PCT / / or PG PUBS #
 Attorney Docket #
 Bar Code #

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CLAIMS

What is claimed is:

1. Use of a biologically active agent in the manufacture of a medicament for treatment of a condition selected from the group consisting of insulin resistance syndrome and diabetes including Type I Diabetes and Type II Diabetes; or for the treatment or reduction in the chance of developing atherosclerosis, arteriosclerosis, obesity, hypertension, hyperlipidemia, fatty liver disease, nephropathy, neuropathy, retinopathy, foot ulceration or cataracts associated with diabetes; or for the treatment of a condition selected from the group consisting of hyperlipidemia, cachexia, and obesity;

wherein the agent is selected from the group consisting of:

4-(4-benzyloxy-3-chlorophenyl)-4-oxobutanoic acid;
Methyl 4-(4-benzyloxy-2-methoxyphenyl)-4-oxobutanoate;
Ethyl 4-(4-cyclohexylmethoxyphenyl)-4-oxobutanoate;
4-(3-chloro-4-cyclopropylmethoxyphenyl)-4-oxobutanoic acid;
Ethyl 3-(4-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(2-benzyloxyphenyl)-3-oxopropanoate;
Methyl 3-(3-(2,6-dichlorobenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(4-(4-chlorobenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(3-(4-methoxybenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(4-methoxybenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(2-methoxybenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(3-methoxybenzyloxy)phenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-chlorophenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-methoxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxy-4-methoxyphenyl)-3-oxopropanoate;

and pharmaceutically acceptable salts thereof.

2. A method for treating a mammalian subject with a condition selected from the group consisting of insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver

disease, cachexia, obesity, atherosclerosis and arteriosclerosis comprising administering to the subject an amount of the biologically active agent effective to treat the condition; wherein the agent is selected from the group consisting of:

4-(4-benzyloxy-3-chlorophenyl)-4-oxobutanoic acid;
Methyl 4-(4-benzyloxy-2-methoxyphenyl)-4-oxobutanoate;
Ethyl 4-(4-cyclohexylmethoxyphenyl)-4-oxobutanoate;
4-(3-chloro-4-cyclopropylmethoxyphenyl)-4-oxobutanoic acid;
Ethyl 3-(4-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(2-benzyloxyphenyl)-3-oxopropanoate;
Methyl 3-(3-(2,6-dichlorobenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(4-(4-chlorobenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(3-(4-methoxybenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(4-methoxybenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(2-methoxybenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(3-methoxybenzyl)oxy)phenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-chlorophenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-methoxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxy-4-methoxyphenyl)-3-oxopropanoate;

and pharmaceutically acceptable salts thereof.

3. The method of claim 2, wherein the agent is administered orally.
4. The method of claim 2, wherein the subject is a human.
5. The method of claim 4, wherein the agent is administered in an amount from one milligram to four hundred milligrams per day.
6. The method of claim 2, wherein the condition is insulin resistance syndrome or Type II Diabetes.
7. The method of claim 2, wherein the condition is Type I Diabetes.

8. The method of claim 2, wherein the treatment reduces a symptom of diabetes or the chances of developing a symptom of diabetes, wherein the symptom is selected from the group consisting of: atherosclerosis, obesity, hypertension, hyperlipidemia, fatty liver disease, nephropathy, neuropathy, retinopathy, foot ulceration and cataracts, associated with diabetes.

9. A pharmaceutical composition for use in the treatment of a condition selected from the group consisting of insulin resistance syndrome, diabetes, hyperlipidemia, fatty liver disease, cachexia, obesity, atherosclerosis, arteriosclerosis and adapted for oral administration, comprising from one milligram to four hundred milligrams of biologically active agent selected from the group consisting of:

4-(4-benzyloxy-3-chlorophenyl)-4-oxobutanoic acid;
Methyl 4-(4-benzyloxy-2-methoxyphenyl)-4-oxobutanoate;
Ethyl 4-(4-cyclohexylmethoxyphenyl)-4-oxobutanoate;
4-(3-chloro-4-cyclopropylmethoxyphenyl)-4-oxobutanoic acid;
Ethyl 3-(4-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxyphenyl)-3-oxopropanoate;
Ethyl 3-(2-benzyloxyphenyl)-3-oxopropanoate;
Methyl 3-(3-(2,6-dichlorobenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(4-(4-chlorobenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(3-(4-methoxybenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(4-methoxybenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(2-methoxybenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(2-(3-methoxybenzyl)phenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-chlorophenyl)-3-oxopropanoate;
Ethyl 3-(4-benzyloxy-3-methoxyphenyl)-3-oxopropanoate;
Ethyl 3-(3-benzyloxy-4-methoxyphenyl)-3-oxopropanoate;

and pharmaceutically acceptable salts thereof.

10. The invention substantially as described above.

L4 1 ANSWERS HCAPLUS COPYRIGHT 2006 ACS on STN
IC ICM A61K
CC 1-10 (Pharmacology)
TI Oxocarboxylic acids and esters thereof for the treatment of metabolic disorders
ST oxocarboxylic acid metabolic disorder treatment; ester oxocarboxylic acid metabolic disorder treatment
IT Antiarteriosclerotics
(antiatherosclerotics; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Kidney, disease
(diabetic nephropathy; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Nerve, disease
(diabetic neuropathy; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Eye, disease
(diabetic retinopathy; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Liver, disease
(fatty; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Ulcer
(foot; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Autoimmune disease
(insulin-dependent diabetes mellitus; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Diabetes mellitus
(insulin-dependent; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Metabolic disorders
(metabolic syndrome X; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Diabetes mellitus
(non-insulin-dependent; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Drug delivery systems
(oral; oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Antiarteriosclerotics
Antidiabetic agents
Antihypertensives
Antiobesity agents
Antiulcer agents
Arteriosclerosis
Atherosclerosis
Cachexia
Cataract
Diabetes mellitus
Drug delivery systems
Human
Hypertension
Hypolipemic agents
Obesity
(oxocarboxylic acids and esters for treatment of metabolic disorders)
IT Hyperlipidemia
RL: BSU (Biological study, unclassified); BIOL (Biological study)

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(oxocarboxylic acids and esters for treatment of metabolic disorders)

IT Foot

(ulcer; oxocarboxylic acids and esters for treatment of metabolic disorders)

IT 13335-57-4 39208-08-7 53090-45-2 60525-32-8 63539-02-6
73083-19-9 74362-70-2 74362-73-5 77513-51-0 102513-61-1
202577-82-0 371251-24-0 373596-81-7 373596-82-8 373596-84-0
387844-34-0

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)

(oxocarboxylic acids and esters for treatment of metabolic disorders)

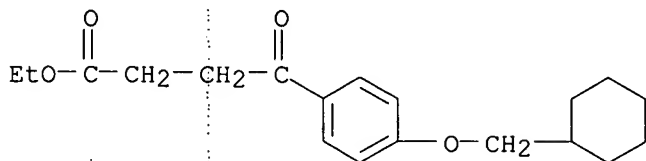
ALL ANSWERS HAVE BEEN SCANNED

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Similar
structures from Inventor

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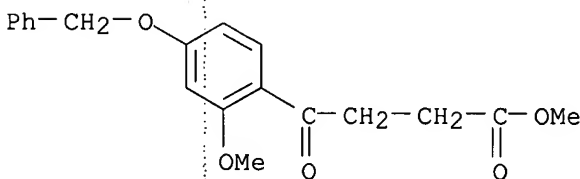
L6 4 ANSWERS REGISTRY COPYRIGHT 2006 ACS on STN
IN *Benzenebutanoic acid, 4-(cyclohexylmethoxy)- γ -oxo-, ethyl ester* (9CI)
MF C19 H26 O4



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

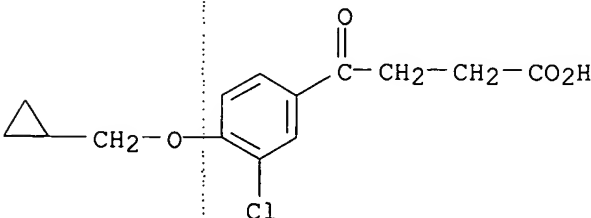
L6 4 ANSWERS REGISTRY COPYRIGHT 2006 ACS on STN
IN *Benzenebutanoic acid, 2-methoxy- γ -oxo-4-(phenylmethoxy)-, methyl ester* (9CI)
MF C19 H20 O5



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L6 4 ANSWERS REGISTRY COPYRIGHT 2006 ACS on STN
IN *Benzenebutanoic acid, 3-chloro-4-(cyclopropylmethoxy)- γ -oxo-* (9CI)
MF C14 H15 Cl O4

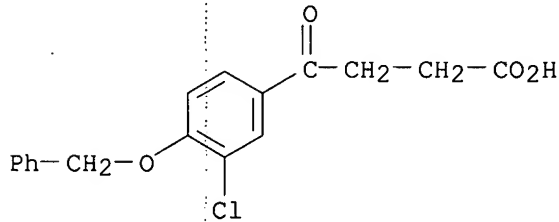


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PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

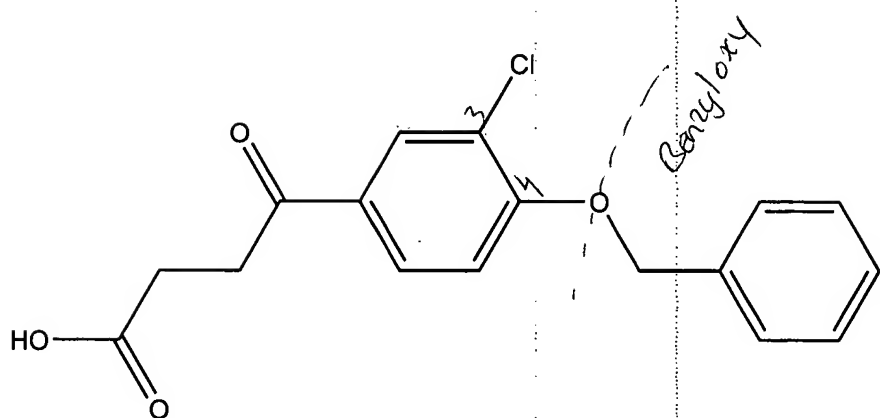
L6 4 ANSWERS REGISTRY COPYRIGHT 2006 ACS on STN
IN *Benzenebutanoic acid, 3-chloro-γ-oxo-4-(phenylmethoxy)-* -
(9CI)
MF C17 H15 Cl O4



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

Chem Draw



4-(4-benzyloxy-3chlorophenyl)-4-oxobutanoic acid